



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6 : C07K 1/02, 1/04, 1/08		A1	(11) International Publication Number: WO 98/56807 (43) International Publication Date: 17 December 1998 (17.12.98)
<p>(21) International Application Number: PCT/US98/12278</p> <p>(22) International Filing Date: 12 June 1998 (12.06.98)</p> <p>(30) Priority Data: 60/049,553 13 June 1997 (13.06.97) US</p> <p>(71) Applicant: GRYPHON SCIENCES [US/US]; Suite 90, 250 East Grand Avenue, South San Francisco, CA 94080-6209 (US).</p> <p>(72) Inventors: CANNE, Lynne; 1103 Crespi Drive, Pacifica, CA 94044 (US). KENT, Stephen, B., H.; 273 Hartford, San Francisco, CA 94114 (US). SIMON, Reyna; 18439 Las Cumbres Road, Los Gatos, CA 95030 (US).</p> <p>(74) Agents: NAKAMURA, Jackie, N.; Cooley Godward LLP, Five Palo Alto Square, 3000 El Camino Reale, Palo Alto, CA 94306-2155 (US) et al.</p>		<p>(81) Designated States: AU, CA, JP, KR, European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE).</p> <p>Published <i>With international search report. Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.</i></p>	
<p>(54) Title: SOLID PHASE NATIVE CHEMICAL LIGATION OF UNPROTECTED OR N-TERMINAL CYSTEINE PROTECTED PEPTIDES IN AQUEOUS SOLUTION</p> <pre> graph TD Handle[HANDLE] --> Resin[COUPLE TO RESIN] Resin --> CoupledHandle[HANDLE-COSR] CoupledHandle --> Cysteine1[LIGATE TO Cys-COS-] Cysteine1 --> Thioester1[CONVERT TO THIOESTER] Thioester1 --> Cysteine2[LIGATE TO Cys-COS-] Cysteine2 --> Thioester2[CONVERT TO THIOESTER] Thioester2 --> Cysteine3[LIGATE TO Cys-COS-] Cysteine3 --> Thioester3[CONVERT TO THIOESTER] Thioester3 --> Cleavage[CLEAVE HANDLE] Cleavage --> FreePeptide[FREE PEPTIDE/PROTEIN] </pre> <p>REPEAT CYCLE AS NEEDED</p>			
<p>(57) Abstract</p> <p>The present invention provides methods, apparatus and kits for synthesizing assembled peptides and proteins on a solid phase with sequential ligation of three or more unprotected peptide segments using chemoselective and mild ligation chemistries in aqueous solution. Also provided are methods of monitoring solid phase sequential ligation reactions using MALDI or electrospray ionization mass spectrometry of reaction products.</p>			

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
AU	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
AZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav Republic of Macedonia	TM	Turkmenistan
BF	Burkina Faso	GR	Greece			TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
BJ	Benin	IE	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
CA	Canada	IT	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NE	Niger	VN	Viet Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	ZW	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's Republic of Korea	NZ	New Zealand		
CM	Cameroon	KR	Republic of Korea	PL	Poland		
CN	China	KZ	Kazakhstan	PT	Portugal		
CU	Cuba	LC	Saint Lucia	RO	Romania		
CZ	Czech Republic	LI	Liechtenstein	RU	Russian Federation		
DE	Germany	LK	Sri Lanka	SD	Sudan		
DK	Denmark	LR	Liberia	SE	Sweden		
EE	Estonia			SG	Singapore		

**SOLID PHASE NATIVE CHEMICAL LIGATION
OF UNPROTECTED OR N-TERMINAL CYSTEINE PROTECTED PEPTIDES
IN AQUEOUS SOLUTION**

5

INTRODUCTION

Background

Existing methods for the chemical synthesis of proteins include stepwise solid phase synthesis, and fragment condensation either in solution or on solid phase. The classic stepwise solid phase synthesis of Merrifield involves covalently linking an amino acid corresponding to the carboxy-terminal amino acid of the desired peptide chain to a solid support and extending the polypeptide chain toward the amino end by stepwise coupling of activated amino acid derivatives having activated carboxyl groups. After completion of the assembly of the fully protected solid phase bound peptide chain, the peptide-solid phase covalent attachment is cleaved by suitable chemistry and the protecting groups removed to give the product polypeptide.

Some disadvantages of the stepwise solid phase synthesis method include: incomplete reaction at the coupling and deprotection steps in each cycle results in formation of solid-phase bound by products. Similarly, side reactions due to imperfections in the chemistry, and or impurities present in the reagents/protected amino acids, all lead to a multiplicity of solid phase bound products at each step of the chain assembly and to the formation of complex product mixtures in the final product. Thus, the longer the peptide chain, the more challenging it is to obtain high-purity well-defined products. Due to the production of complex mixtures, the stepwise solid phase synthesis approach has size limitations. In general, well-defined polypeptides of 100 amino acid residues or more are not routinely prepared via stepwise solid phase synthesis. Synthesis of proteins and large polypeptides by this route is a time-consuming and laborious task.

The solid phase fragment condensation approach (also known as segment condensation) was designed to overcome the difficulties in obtaining long polypeptides via the solid phase stepwise synthesis method. The segment condensation method involves preparation of several peptide segments by the solid phase stepwise method, followed by cleavage from the solid phase and purification of these maximally protected segments. The protected segments are condensed one-by-one to the first segment, which is bound to the solid phase.

Often, technical difficulties are encountered in many of the steps of solid phase segment condensation. See E. Atherton, et al., "Solid Phase Fragment Condensation - The Problems," in *Innovation and Perspectives in Solid Phase Synthesis* 11-25 (R. Epion, et al. 5 1990). For example, the use of protecting groups on segments to block undesired ligating reactions can frequently render the protected segments sparingly soluble, interfering in efficient activation of the carboxyl group. Limited solubility of protected segments also can interfere with purification of protected segments. See K. Akaji et al., *Chem. Pharm. Bull.(Tokyo)* 33:184-102 (1985). Protected segments are difficult to characterize with 10 respect to purity, covalent structure, and are not amenable to high resolution analytical ESMS (electrospray mass spectrometry) (based on charge). Racemization of the C-terminal residue of each activated peptide segment is also a problem, except if ligation is performed at Glycine residues. Moreover, cleavage of the fully assembled, solid-phase bound polypeptide from the solid phase and removal of the protecting groups frequently can require harsh chemical 15 procedures and long reaction times that result in degradation of the fully assembled polypeptide.

Segment condensation can be done in solution rather than on solid phase. See H. Muramatsu et al., *Biochem. and Biophys. Res. Commun.* 203(2):1131-1139 (1994). However, segment condensation in solution requires purification of segments prior to ligation as well as 20 use of protecting groups on a range of different side chain functional groups to prevent multiple undesired side reactions. Moreover, the ligation in solution does not permit easy purification and wash steps afforded by solid phase ligations. Furthermore, the limitations with respect to solubility of protected peptide segments and protected peptide intermediate reaction products are exacerbated.

25 Chemical ligating of minimally protected peptide segments has been explored in order to overcome the solubility problems frequently encountered with maximally protected peptide segments. See Cheng, et al., *Chemical Synthesis of Human 9-endorphin(1-27) Analogs by Peptide Segment Coupling*. *Int. J. Pept. Protein Res.* 38:70-78 (1991); J. Blake, *Total Synthesis of S-Carbamoylmethyl Bovine Apocytochrome c by Segment Coupling*, *Int. 30 J. Pept. Protein Res.* 27:191-200 (1986); and H. Hojo et al., *Protein Synthesis using S-Alkyl Thioester of Partially Protected Peptide Segments, Synthesis of DNA-Binding Protein of Bacillus stearothermophilus*, *Bull. Chem. Soc. Jpn.* 65:3055-3063 (1992). However, this method still requires the use of protecting groups on all Lysine side chain amino groups, selective N- α protection of one or more segments, and laborious purification steps, involving 35 purification, reprotection, and repurification.

The use of multiply protected peptide segments is incompatible with the overall scheme of engineering proteins using peptides produced by means of recombinant DNA expression as a source. Protected peptide segment methods are labor-intensive, and the protected peptide segments have unpredictable handling properties, partly due to the 5 solubility and ligating difficulties of protected peptide segments. Often, large protected peptide segments are minimally soluble in even the most powerful polar aprotic solvents such as dimethylsulfoxide (DMSO) and dimethylformamide (DMF). The problem of insolubility in protected peptide segments has been addressed with limited success in several ways, including the use of (1) partial protecting group strategy which masks all side chains except 10 those of Ser, Thr, and Tyr; (2) minimal protecting group strategy that masks only thiol and amino side chains; and (3) using reversible protection of a backbone amide moiety to prevent aggregation/insolubility. Protecting groups used in the latter approach alter peptide conformations. Use of backbone protecting groups is not yet straightforward or predictable and requires significant experimentation for each target polypeptide chain.

15 There are a number of techniques for ligating unprotected peptide segments via unnatural backbone linkages. In contrast, there are few methods for achieving a "native chemical ligation." A "native chemical ligation" is the chemoselective reaction of unprotected or N-terminal Cysteine protected peptide segments with another unprotected peptide segment resulting in the formation of a ligated peptide with an amide bond at the 20 ligation site. The fully assembled target polypeptides of the invention comprise one, two or more native chemical ligation sites.

Accordingly, there is a need in the art for rapid methods of synthesizing assembled polypeptides via chemical ligation of two or more unprotected peptide segments using a solid support, with improved yields and facilitated handling of intermediate products.

25 The present invention makes possible, inter alia, the rapid solid-phase synthesis of large polypeptides with a natural peptide backbone via native chemical ligation of two or more unprotected peptide segments where none of the reactive functionalities on the peptide segments need to be temporarily masked by a protecting group. The present invention accomplishes for the first time, solid phase sequential chemical ligation of peptide segments 30 in an N-terminus to C-terminus direction, with the first solid phase-bound unprotected peptide segment bearing a C-terminal α -thioester that reacts with another unprotected peptide segment containing an N-terminal Cysteine and a C-terminal thioacid.

Other embodiments of the invention also permit solid-phase native chemical ligation in the C- to N-terminus direction, with temporary protection of N-terminal cysteine residues 35 on an incoming (second) peptide segment. Those of ordinary skill in the art will readily

appreciate that the invention may also include the use of nonnative chemical ligation to sequentially ligate peptide segments via unnatural linkages on a solid phase. Alternatively, the invention may include the use of native chemical ligation of peptide segments wherein said peptide segments comprise one or more unnatural backbone linkages.

5

References

- Matthys J. Janssen, "Thiolo, Thiono, and Dithio Acids and Esters," Chptr. 15 of The Chemistry of Carboxylic Acids and Their Esters (1969).
- Schnolzer et al., Science 256:221-225 (1992)
- 10 Rose et al. J. Am Chem. Soc. 116:30-34 (1994)
- Liu et al., Proc. Natl. Acad. Sci. USA 91:6584-6588 (1994).
- Dawson et al. Science 266:77-779 (1994).
- PCT/US95/05668, WO 96/34878
- Sakakibara S., Biopolymers (Peptide Science), 37:17-28 (1995).
- 15 Tam et al., PNAS USA, 92:12485-12489 (1995).

SUMMARY OF THE INVENTION

The present invention provides, inter alia, novel methods of producing large polypeptides by native chemical ligation of peptide segments in aqueous solution to an unprotected solid phase bound peptide without need for protecting groups on the peptide segments, or, with temporary protection of the N-terminal cysteine of incoming peptide segments. Among the many advantages of this embodiment of the invention are: ease of purification of the intermediate and final products; faster ligation reactions; rapid synthesis of large polypeptides with a natural peptide backbone; ease of ligating reactions due to the lack of protecting groups and resultant enhanced solubility of peptide segments in aqueous or mixed aqueous/organic solutions; chemoselective ligation due to the lack of reactivity of the thioester moiety with other functional groups present in both reactive peptide segments to form stable co-products, resulting in a purer final product without side reactions; adaptability to monitoring on the solid phase via MALDI mass spectrometry or ESI MS (electrospray ionization mass spectrometry); decreased racemization due to the use of mild activation using a thioester and the avoidance of elevated pHs; the polypeptide product is obtained directly in unprotected form; and adaptability to automation and combinatorial techniques.

A significant advantage of the solid phase ligations over solution ligations is that the solid phase ligation methods do not require arduous HPLC (high pressure liquid chromatography) purification and lyophilization steps after each ligating reaction, whereas

ligations in solution do. Thus, the solid phase ligations eliminate many time-consuming purification steps that decrease the recovery of final product. Instead, the solid phase sequential ligation methods here described only require a single HPLC purification and lyophilization step after the final unprotected peptide segment has been ligated and the 5 assembled peptide is cleaved from the solid phase. The elimination of these time-consuming purification steps allows for faster synthesis of the final product, i.e. the assembled peptide, than would the analogous route in solution. Ready purification of the desired solid phase-bound product from soluble coproducts presents a tremendous advance in terms of the yield of the ultimate assembled polypeptide.

10 Another advantage of solid phase ligations is that they permit higher concentrations of reactants which leads to faster reaction rates. For example, by using an excess at high concentration of the incoming peptide segment as compared to the solid phase-bound peptide, reactions can reach completion much faster. The excess peptide segment can readily be washed off the solid phase after the ligation reaction is complete. Increased yields of final 15 product can be accomplished by increasing concentrations of peptide segments. For example, the solid phase-bound polypeptide can be dried out on the solid-phase and ressolvated in ligation solution. Alternatively, the solid phase-bound peptide can be washed with a solution of incoming peptide segments at high concentration.

Other advantages of the present invention are that it allows for synthesis of much 20 larger peptides and proteins than are presently attainable by conventional methods, it is amenable to automation, and the use of high resin loadings allow for easy scale up. Moreover, ligation in the N- to C-terminal direction permits the use of crude peptide segments without need for purification or lyophilization, since termination products formed during stepwise solid phase synthesis of the peptide segments will be unreactive with the 25 solid phase-bound peptide.

In one embodiment, the invention comprises a method of producing an assembled peptide having a native peptide backbone by ligating peptide segments in the N- to C-terminal direction, comprising: a) covalently binding an unprotected first peptide segment to a solid phase via a linker comprising a cleavable moiety, wherein said cleavable moiety is 30 stable under ligation conditions and said unprotected first peptide segment is bound to said cleavable moiety at its N-terminus and has an α -thioester at its C-terminus; b) optionally introducing a second unprotected peptide segment, wherein said second segment comprises a cysteine residue at its N-terminus and a thioacid at its C-terminus, under conditions suitable to permit ligation between said first unprotected peptide segment and said second unprotected 35 peptide segment to form a natively ligated peptide bound to said solid phase, wherein said

solid phase-bound peptide comprises a thioacid at its C-terminus, and subsequently converting said solid phase-bound peptide thioacid to a thioester; (c) optionally repeating step (b) with additional unprotected peptide segments; (d) introducing a final unprotected peptide segment, comprising a cysteine residue at its N-terminus, under conditions suitable to permit 5 ligation between said solid phase-bound peptide and said final unprotected peptide segment. In a preferred embodiment, the cleavable moiety is cleaved to release the solid phase-bound peptide in the form of the assembled peptide. In another preferred embodiment, cleavable moiety is a cleavable linker capable of being cleaved for purposes of monitoring the sequential ligation reactions. In another embodiment, the first unprotected peptide segment is 10 added as a peptide- α COSH thioacid and subsequently converted to a thioester.

The sequential ligation in the N- to C-terminus direction is a surprisingly effective and elegant means of obtaining chemoselective ligation of unprotected peptide segments without racemization. Before the present invention, sequential ligations were not conducted in the N- to C-terminal direction due to concerns regarding racemization at the α COX at the 15 C-terminus of the peptide (peptide- α COX). Using the present invention, the α COSH at the C-terminus of the peptide segment is mildly activated to a thioester and the ligating reaction is carried out in the absence of base, in an aqueous buffered solution, resulting in mild conditions that do not generate racemic mixtures.

The methods of the invention can be used for native chemical ligation of peptide 20 segments produced by stepwise solid phase synthesis. The last peptide segment to be added at the C-terminal end of the last solid phase-bound peptide in the reaction scheme may be a recombinantly expressed peptide having an N-terminal Cysteine residue (Cys-recombinant peptide). The thioacid moiety, which is activated to a thioester moiety, can be placed anywhere a native chemical ligation is desired, including on a side chain. Thus, the sequential 25 ligations of the invention are not limited to linearly assembled peptides.

In another embodiment, there is the use of unprotected peptide segment middle pieces each having an N-terminal cysteine residue that participate in native chemical ligation.

In another embodiment, the invention comprises a method of producing an assembled peptide having a native peptide backbone by ligating peptide segments in the C- to N- 30 terminal direction, comprising: a) covalently binding an unprotected first peptide segment to a solid phase via a cleavable handle comprising a cleavable moiety, wherein said cleavable moiety is stable under ligation conditions and said unprotected first peptide segment is bound to said cleavable moiety at its C-terminus and has a Cysteine at its N-terminus; b) introducing a second peptide segment, wherein said second segment comprises a cysteine residue at its N- 35 terminus and an alpha-thioester at its C-terminus, and wherein said second peptide segment

has a protecting group bound to its N-terminal cysteine residue, under conditions suitable to permit ligation between said first peptide segment and said second N-terminally protected peptide segment to form a natively ligated peptide bound to said solid phase, wherein said solid phase-bound peptide comprises a protecting group bound to an N-terminal cysteine; c) 5 removing said protecting group from solid phase-bound peptide; (d) optionally repeating steps b) and c) with additional peptide segments comprising an N-terminal Cysteine and a C-terminal alpha thioester, wherein said additional peptide segments have a protecting group bound to their N-terminal cysteine residue (e) introducing a final peptide segment, comprising an alpha-thioester at its C-terminus, providing that if said final peptide segment 10 comprises an N-terminal Cysteine, said N-terminal Cysteine is protected by a protecting group, wherein said introducing occurs under conditions suitable to permit ligation between said solid phase-bound peptide and said final peptide segment; and (e) optionally removing said protecting group from the N-terminal cysteine of said solid phase-bound peptide.

In another embodiment, the invention comprises a method of producing an assembled 15 peptide having a native peptide backbone by ligating peptide segments in the C- to N-terminal direction, comprising: a) covalently binding an unprotected first peptide segment to a solid phase via a cleavable handle comprising a cleavable moiety, wherein said cleavable moiety is stable under ligation conditions and said unprotected first peptide segment is bound to said cleavable moiety at its C-terminus and has a Cysteine at its N-terminus; b) optionally 20 introducing a second peptide segment, wherein said second segment comprises a cysteine residue at its N-terminus and an alpha-thioester at its C-terminus, and wherein said second peptide segment has a protecting group bound to its N-terminal cysteine residue, under conditions suitable to permit ligation between said first peptide segment and said second N-terminally protected peptide segment to form a natively ligated peptide bound to said solid 25 phase, wherein said solid phase-bound peptide comprises a protecting group bound to an N-terminal cysteine, and subsequently removing said protecting group from solid phase-bound peptide; (c) optionally repeating step (b) with additional peptide segments comprising an N-terminal Cysteine and a C-terminal alpha thioester, wherein said additional peptide segments have a protecting group bound to their N-terminal cysteine residue; (d) introducing a final 30 peptide segment, comprising an alpha-thioester at its C-terminus, providing that if said final peptide segment comprises an N-terminal Cysteine, said N-terminal Cysteine is protected by a protecting group, wherein said introducing occurs under conditions suitable to permit ligation between said solid phase-bound peptide and said final peptide segment; and (e) optionally removing said protecting group from the N-terminal cysteine of said solid phase- 35 bound peptide.

In yet another embodiment, there is the solid phase sequential ligation of peptide segments in either or both directions, using a cleavable linker to monitor the ligation reactions via mass spectrometry and to purify the assembled peptide from the solid phase.

Another embodiment is a method of bidirectional solid phase native chemical ligation,
5 comprising providing a first peptide segment bound to a solid support via one of its internal amino acid residues, wherein said first peptide segment comprises an N-terminal Cysteine and a C-terminal thioester, and ligating a second peptide segment to either terminus.

In another embodiment, there is provided a kit comprising an unprotected peptide segment, covalently bound via an internal amino acid side chain functional group to a
10 cleavable handle, wherein said cleavable handle is linked to a solid phase via a chemoselective functional group complementary to a chemoselective functional group on the solid phase. Said kit can be used for solid phase chemical ligation of unprotected or N-terminal cysteine-protected peptide segments to the solid phase-bound peptide. A preferred example of such a cleavable handle is

15 a functionalized cleavable handle, X-aminoethylsulfonylethyoxy carbonyl (wherein X=CH₃COCH₂CH₂CH₂CONHCH₂-MSC- or X= AOA-NHCH₂-MSC-. (AOA = aminoxyacetal).

In another embodiment, there are methods of using bromoacetic acid or iodoacetic acid to convert a peptide segment thioacid (peptide- α COSH) to a thioester (peptide- α COSR),
20 on a solid phase.

In yet another embodiment, there is provided a method of monitoring the solid phase sequential ligation process on the solid phase via MALDI or ESI mass spectrometry, using cleavable linkers. Monitoring via ESI MS can also be accomplished using a TFA-cleavable linker or, when MALDI is the mass spectrometric method used, a photocleavable linker may
25 preferably be used.

In a further embodiment, there are provided novel methods of preparing modular large peptide or protein libraries using combinations of the aspects of the invention described herein. Particularly useful are the methods of solid phase sequential ligation of peptide segments to rapidly synthesize multiple analogs of known proteins or polypeptides.

30 Kits and apparatus for assembling polypeptides and polypeptide libraries by the processes described herein are also provided.

One of skill in the art will readily appreciate that each of the embodiments of the invention can be combined with other embodiments to obtain a wide range of useful inventions.

BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 is a schematic diagram of a solid phase native chemical ligation scheme, in the N- to C-terminus direction. In one embodiment, the linker is an MSC handle, which is cleavable yet stable under ligation conditions. In another embodiment, the unprotected first 5 peptide segment is covalently bound to a solid phase (resin) via an aminoxy-ketone linkage.

FIG. 2A illustrates the stability of a 13-residue peptide- α -COSH with a Cysteine residue at the N-terminus under ligation conditions. The HPLC chromatogram shows that only a small percentage of the peptide cyclized or formed larger aggregates, even after overnight storage under ligation conditions.

10 **FIG. 2B** illustrates the stability of the same 13-residue peptide- α COSH in the presence of a thioester peptide having a molecular weight of 1230.2. The HPLC chromatogram shows that the Cys- α -COSH peptide is adequately stable to use in ligation without significant reaction with itself. Furthermore, such byproducts as are formed in small proportion by reaction of the 13-residue peptide- α COSH (having an N-terminal cysteine) 15 with itself are unreactive with a resin-bound peptide α -COSR and are readily removed by simple filtration and washing.

20 **FIG. 3A, 3B and 3C** show HPLC chromatograms of the effect of hydrazine on the removal of the MSC handle from a peptide having an N-terminal Cysteine residue. The peak correlating with the mass of 1708.2 represents the desired peptide with the MSC handle 25 removed. The peak corresponding to the mass of 1814.5 represents a reactive side product formed upon cleavage that can react with the desired peptide without the MSC handle. **FIG. 3A** shows a fairly large peak at the 1814.5 mw when an aliquot of the peptide was placed in 6M guanidine• HCl, 0.1 M NaPi, pH 7.5, then diluted into 1 N NaOH for 2 min., then quenched with 1 N HCl. **FIG. 3B** is an HPLC chromatogram of the resulting product when 30 the conditions of **FIG. 3A** are repeated with the inclusion of 50 mM hydrazine in the 6 M guanidine• HCl solution. **FIG. 3C** is an HPLC chromatogram of the resulting product when the conditions of **FIG. 3A** are repeated with 200 mM hydrazine in the 6 M guanidine• HCl solution. Hydrazine scavenges the side product, resulting in a purer product.

35 **FIG. 4** is an HPLC chromatogram of the removal of a cleavable MSC handle from a peptide that does not have an N-terminal Cysteine residue, but rather an N-terminal Leucine residue and a Cysteine residue in its approximate center. The molecular weight of the peptide with the MSC handle is 4022.4 and without the MSC handle, 3745.1. An aliquot of the peptide in 6M guanidine• HCl, 0.1M NaAc, pH 4.6 was diluted into 6M guanidine• HCl, 0.1M NaAc, pH 14 for 2 min., quenched with 6M guanidine• HCl, 0.1M NaAc, pH 2.0. The HPLC shows that an internal reaction with the side product still occurs, to form the peak

having a mw of 3979.7 (corresponding to the modification by the LEV-NHCH₂- handle), but that the extent of the reaction is less than that occurring with a peptide having an N-terminal Cysteine.

FIG. 5A is a reaction scheme showing the preparation of the PEGA resin used as the 5 solid support in N- to C-terminal sequential ligations. Steps A and B1 are optional steps to produce a photolabile linker for use with MALDI analysis of the resin samples.

FIG. 5B is a diagram illustrating a generalized scheme for preparing a solid phase (resin) for use in the solid phase sequential ligations of the invention. Structure 1 is a cleavable linker useful for monitoring the progress of coupling and ligation reactions by mass 10 spectrometry. For example, a photo-cleavable linker can be used for on-resin monitoring by MALDI MS, whereas a TFA cleavable linker can be used for monitoring by electrospray MS. Once structure 1 is coupled to the resin, the protecting group (PG) is removed and a functional moiety (structure 3) capable of chemoselective reaction with the first peptide segment, is added to the resin. Once 3 is coupled to the resin, the protecting group is 15 removed to give structure 4, which is ready for chemoselective reaction with structure 5, a peptide modified with a cleavable handle and a functional group capable of reaction with the now modified resin (4). Once all subsequent ligations are complete, the "cleavable handle" is cleaved to release the full length peptide (assembled peptide) from the solid phase.

FIG. 6 is a reaction scheme illustrating the derivatization of Peptide Segment 1 (the 20 N-terminal peptide segment).

FIG. 7A and 7B are HPLC chromatograms of the coupling of a first unprotected peptide segment (1) to the solid support, in this example, an AOA-functionalized resin (PEGA). FIG. 7A is an HPLC of the peptide solution as added to the resin. FIG 7B is an HPLC of the supernatant after reaction of the peptide with a molar excess of the resin 25 overnight. A significant amount of the peptide has been removed from the supernatant, indicating that it has been bound to the resin after the overnight reaction.

FIG. 8A and 8B are HPLC chromatograms of the same experiments reflected in FIG. 7A and 7B, except with Isco resin beads as the solid phase.

FIG. 9A, 9B, and 9C are analyses of the products after step 1 of this figure, binding 30 of the first unprotected peptide segment to the solid phase. **FIG. 9A** is an analytical HPLC chromatogram of the (base plus hydrazine) cleavage of the resin-bound peptide. **FIG. 9B** is a MALDI mass spectrum of the resin, showing a peak corresponding to (1), the resin-bound peptide. **FIG. 9C** is a MALDI mass spectrum after base cleavage of the linker, showing the lack of a peak corresponding to (1), and showing that no peptide is sticking to the solid phase 35 (resin).

FIG. 10A, 10B, and 10C are analyses of the products after step 3 of this figure, i.e., ligating of the second unprotected peptide segment (2) to the resin-bound peptide (1). **FIG. 10A** is an analytical HPLC of the product, resin-bound peptide intermediate, showing a large peak with mass of (1) + (2). **FIG. 10B** is a MALDI mass spectrum of the resin before 5 cleavage of the linker, and **FIG. 10C** is a MALDI mass spectrum of the resin after base cleavage of the linker.

FIG. 11 is an HPLC chromatograph of the desalted, lyophilized peptide product (1+2+3 of Table 1) after 2 sequential ligations on a solid phase (Isco resin) in the N- to C-terminal direction. The tallest peak corresponds to the crude, lyophilized product, indicating 10 approximately 36% yield.

FIG. 12A and 12B are ESI MS (electrospray ionization mass spectra) of the main peak corresponding to the assembled peptide (1+2+3 of Table 1). **FIG. 12B** is a reconstructed display of the mass spectrum of FIG. 12A, showing the mass of the product ligated peptide.

15 **FIG. 13** is an HPLC chromatogram of the desalted, lyophilized peptide (1+2+3) after base cleavage of the linker to remove the assembled peptide from the solid phase (PEGA resin).

FIG. 14A and 14B are electrospray ionization mass spectra of the 7434 mass peak, wherein FIG. 14B is a reconstruction of the mass spectrum of FIG. 14A.

20 **FIG. 15A, 15B and 15C** are 3 HPLC chromatograms illustrating that the solid support technique can be used for both purification and ligation. Figs. 15A and 15B show solution processing of a crude peptide before and after removal of DNP groups, respectively. Both HPLCs show a crude mixture of peptides. Fig. 15C is an HPLC chromatogram of the same peptide solution shown in FIG. 15A, after coupling to a solid support, removal of DNP 25 groups and base cleavage from the solid phase, resulting in a significantly purer assembled peptide product.

FIG. 16A and 16B illustrate the reaction scheme for synthesis of MIF(1-115) via solid phase sequential native ligations in the N-terminal to C-terminal direction.

30 **FIG. 17A** is a reaction scheme for the modification of the N-terminal peptide segment. **FIG. 17B** is a diagram illustrating the modification of the aqueous-compatible solid phase in preparation for coupling the first unprotected peptide segment.

FIG. 18A is a reaction scheme for the coupling of N-terminal modified MIF(1-59) to a solid phase. **FIG. 18B** is an HPLC chromatogram of the released peptide after base cleavage, having an expected mass of 6271 Da. **FIG. 18C and 18D** are electrospray mass

spectra of the main component of the released peptide after cleavage of the cleavable handle.

FIG. 18D is a reconstruction of **FIG. 18C**.

FIG. 19A is a diagram of the ligation step to form resin-bound MIF(1-80). **FIG. 19B** is an HPLC chromatogram of the products after cleavage of the cleavable handle. **FIG. 19C** and **19D** are mass spectra of the main components of the released peptide after base cleavage, having an expected mass of 8502 Da. **FIG. 19D** is a reconstructed display of the mass spectrum of **FIG. 19C**.

FIG. 20A is a diagram of the ligation step to form resin-bound MIF(1-115). **FIG. 20B** is an HPLC chromatogram of the products after cleavage of the cleavable handle. **FIG. 20C** and **FIG 20D** are mass spectra of the released products after base cleavage, having an expected mass of 12450 Da.

FIG. 21 is a schematic diagram of solid phase ligations in the C- to N-terminus direction. The “resin” represents a solid phase. The triangle and its sideways M-shaped partner are complementary functional groups that chemoselectively form a covalent bond. The “handle” is a cleavable handle that can be cleaved to remove the assembled peptide product from the solid phase. The undulating lines comprise amino acid residues of peptide segments. The “PG” represents a protecting group, which can be placed either on a side chain thiol or on the α -amino group of the N-terminal cysteine. Steps 2 and 3 can be repeated, as indicated by the arrow marked 4, for additional peptide segments. Also, a cleavable linker for purposes of monitoring the coupling and ligating reactions can be added between the “handle” and the “resin.”

FIG. 22 is a reaction scheme for solid phase sequential ligation in the C- to N-terminal direction of PLA2G5.

FIG. 23 is a reaction scheme for synthesizing a Cam ester derivative for solid phase sequential ligation in the C- to N-terminal direction.

FIG. 24 is a reaction scheme for synthesizing the C-terminal peptide segment for solid phase sequential ligation in the C- to N-terminal direction.

FIG. 25A, B, and C is a diagram of a scheme for synthesizing an assembled polypeptide via bidirectional solid phase sequential ligation of two or more peptide segments.

30

FIG 26 A and B are HPLC chromatographs following the solid phase solid phase native chemical ligation of 3 peptide segments in the N- to C- terminal direction, resulting in the assembled peptide, C5a 1-74.

FIG. 27 is a reaction scheme for synthesis of a C-terminal peptide segment for use in the solid phase native chemical ligations described herein, using a CAM ester cleavable handle to remove the synthesized peptide segment from the solid phase.

FIG. 28A and B are HPLC chromatographs and reconstructed ESI MS of the assembled peptide resulting from solid phase sequential ligation of 3 peptide segments: peptide segment 1 (SEQ ID NO: 2)(CADRKNILA), peptide segment 2 (SEQ ID NO: 3)(CYGRLEEKG) and peptide segment 3 (SEQ ID NO: 4)(ALTKYGFYG) on solid phase in the C- to N-terminal direction, using Fmoc protecting groups.

FIG. 29A and B are an HPLC chromatograph and ESI MS, respectively, of the final ligation product, i.e. the first ligation product ligated to the third peptide segment (ALTKYGFYG), resulting from solid phase sequential ligation of 3 peptide segments in the C- to N-terminal direction, using ACM as the protecting group.

FIG. 30A-H are HPLC chromatographs and reconstructed ESI MS of the steps of synthesizing Phospholipase A2 Group 5, a 118 residue protein, using solid phase sequential native chemical ligation of four peptide segments in the C- to N-terminal direction. The first peptide segment is PLA2G5 88-118; the second is PLA2G5 59-87, the third is PLA2G5 26-58, and the fourth is PLA2G5 1-25. **FIG 32A and B** are an HPLC chromatograph and reconstructed ESI MS of the first peptide segment, respectively. **FIG 32C and D** are an HPLC chromatograph and reconstructed ESI MS, respectively, of the ligation product of the first and second peptide segments (PLA2G5 59-118). **FIG 32E and F** are an HPLC chromatograph and reconstructed ESI MS, respectively, of PLA2G5 26-118, the ligation product of PLA2G5 59-118 and PLA2G5 26-58 (the third peptide segment). **FIG 32G and H** are HPLC chromatograph and reconstructed ESI MS, respectively, of PLA2G5 1-118, the assembled polypeptide.

25

DESCRIPTION OF SPECIFIC EMBODIMENTS

Terminology

Amino acids: Amino acids include the 20 genetically coded amino acids, rare or unusual amino acids that are found in nature, and any of the non-naturally occurring and modified amino acids.

Aqueous solution: solutions containing water, including up to 8M urea in water, up to 6M guanidine• HCl in water, up to 60% acetonitrile in water.

Assembled Peptide: the final product of a solid phase sequential or bidirectional ligation, after cleavage of the cleavable handle. The assembled peptide comprises at least

two separate peptide segments sequentially ligated on a solid phase. The assembled peptide may or may not have biological activity.

Cleavable Handle: A cleavable moiety that is capable of being selectively cleaved to release the assembled peptide from the solid phase. The cleavable handle must be capable of 5 resisting cleavage under conditions suitable for coupling, activating, deprotecting, ligating, washing, and other steps involved in the formation of an assembled peptide. The cleavable handle must also be stable to conditions used to produce the first peptide segment that is capable of being bound to a solid phase, including, for example, stepwise solid phase peptide synthesis. The cleavable handle preferably is located directly adjacent to the first peptide 10 segment such that upon cleavage of the cleavable handle, the desired assembled peptide is released from the solid phase. The cleavable handle may be selected from any of the variety of cleavable handles used by those in the field. See, e.g., L. Canne et al., Tetrahedron Letters, 38(19):3361-3364 (1997); Ball et al., J. Pept. Sci, 1:288-294 (1995); Funakoshi et al, PNAS USA, 88:6981-6985 (1991); Funakoshi et al., J. Chromatog. 638:21-27 (1995); Garcia- 15 Echeverria et al., J. Chem. Soc., Chem. Commun., 779-780 (1995). A preferred cleavable handle is Boc-HN-CH₂-CH₂-SO₂-CH₂-CH₂-O-CO-ONp (Boc-HNCH₂-MSC-) or a functionalized cleavable handle, X-aminoethylsulfonyl ethyloxycarbonyl (wherein X=CH₃COCH₂CH₂CH₂CONHCH₂-MSC- or X= AOA-NHCH₂-MSC-). (AOA = aminoxyacetal). Another preferred cleavable handle is a CAM ester. See Ceccato, M.L. et 20 al., Tetrahedron Lett. 31:6189-6192 (1990).

Cleavable Linker: A cleavable moiety that is capable of being selectively cleaved to monitor the solid phase sequential ligation using mass spectrometry of small samples of the reaction mixture at any point during the ligation procedure, i.e. after ligating of the second peptide segment, after ligating of the third peptide segment, and so forth. The cleavable 25 linker must be stable under coupling and ligating conditions, deprotecting conditions (if needed), and washing conditions. Preferred cleavable linkers include photolabile linkers and TFA-labile linkers.

Coupling: Chemoselective reactions involving covalent binding of a first peptide segment to a solid phase.

30 Ligating: Chemoselective reactions involving covalent binding of a peptide segment to a solid phase-bound peptide.

Linker: A covalent linkage linking various moieties. For example, a linker may link a first peptide segment and a solid support, and such a linker may optionally comprises any number of moieties, including a cleavable handle, a cleavable linker, complementary

functional groups capable of chemoselectively forming a covalent bond (e.g., amino-oxy and ketone to form an oxime).

Peptide: A polymer of at least two monomers, wherein the monomers are amino acids, sometimes referred to as amino acid residues, which are joined together via an amide bond. For purposes of this invention, the terms "peptide," "polypeptide," and "protein," are largely interchangeable as all three types can be made via the methods described herein. Peptides are alternatively referred to as polypeptides. Amino acids include the L and D isoforms of chiral amino acids.

Peptide segment: A peptide or polypeptide, having either a completely native amide backbone or an unnatural backbone or a mixture thereof, ranging in size from 2 to 1000 amino acid residues, preferably from 2-99 amino acid residues, more preferably from 10-60 amino acid residues, and most preferably from 20-40 amino acid residues. Each peptide segment can comprise native amide bonds or any of the known unnatural peptide backbones or a mixture thereof. Each peptide segment can be prepared by any known synthetic methods, including solution synthesis, stepwise solid phase synthesis, segment condensation, and convergent condensation. The final peptide segment to be added to form the assembled peptide product can be recombinantly expressed.

Protecting Group: A chemical moiety capable of protecting a functional group from reacting with another functional group, and removable without damage to the formed amino acid or peptide.

Sequential ligation: ligating three or more peptide segments together in order from C-terminus to N-terminus or from the N-terminus to C-terminus, depending on the directionality chosen, to obtain an assembled peptide product. The directionality of the sequential ligations will always start from the solid phase-bound first peptide segment to the last peptide segment to be added to form the assembled peptide product.

Solid Phase: A material having a surface and which is substantially insoluble when exposed to organic or aqueous solutions used for coupling, deprotecting, and cleavage reactions. Examples of solid phase materials include glass, polymers and resins, including polyacrylamide, PEG, polystyrene PEG-A, PEG-polystyrene, macroporous, POROSTM, cellulose, reconstituted cellulose (e.g. Perloza), nitrocellulose, nylon membranes, controlled-pore glass beads, acrylamide gels, polystyrene, activated dextran, agarose, polyethylene, functionalized plastics, glass, silicon, aluminum, steel, iron, copper, nickel and gold. Such materials may be in the form of a plate, sheet, petri dish, beads, pellets, disks, or other convenient forms. Sheets of cellulose can be used as a solid phase in the present invention to accomplish spot ligation in a spatially addressable array. Many of the examples and

embodiments described herein refer to resins, which are a type of solid phase, and one of ordinary skill in the art would understand that such examples are not meant to be limited to resins, but to solid phases in general. The terms solid phase and solid support are used herein interchangeably.

- 5 Solid Phase-bound Peptide: a solid phase-bound peptide comprises at least one peptide segment bound to a solid phase via any variety of cleavable linkers, handles or moieties. A solid phase-bound peptide can include any of the intermediate peptide products of the sequential ligation reactions, including the final solid-phase bound peptide produced after the final peptide segment is ligated to the penultimate solid phase-bound peptide.
- 10 Thioacid: An ionizable thioacid moiety, represented by either -COSH or -COS⁻, often referring to a peptide thioacid, represented by "peptide α-COSH" or "peptide α-COS⁻."

Thioester: A moiety represented by -COSR, often connected to a peptide. For example, a peptide thioester may be represented as "peptide α-COSR". The R group may be
15 any number of groups, including 1-15 C functionalized alkyl, straight or branched, 1-15 C aromatic structures, 1-4 amino acids or derivatives thereof, preferably wherein the R group is selected such that the peptide-alpha-COSR is an activated thioester. In a preferred embodiment, R = -CH₃-Ø, -Ø. The term "thioester" is commonly used, but the true IUPAC term is "thiol ester." See Matthys J. Janssen, *supra*

20

I. SOLID PHASE SEQUENTIAL NATIVE LIGATION OF UNPROTECTED PEPTIDE SEGMENTS IN THE N- TO C-TERMINAL DIRECTION

There have been few reports of proteins synthesized by sequential, multiple ligations of three or more unprotected peptide segments. Such sequential ligations of free peptide
25 segments in solution consequently require a purification (e.g. HPLC) after each ligation and typically require temporary protection of one of the functionalities of the middle segments.

One aspect of the present invention is a solid phase sequential ligation technique which avoids the need for multiple purifications and the need to temporarily protect the middle peptide segments. This strategy employs (1) the modification of the N-terminal
30 peptide segment with a cleavable handle functionalized with a group capable of chemoselective reaction with the solid support and (2) sequential native chemical ligations of unprotected peptide segments in an N- to C-terminal direction. Native chemical ligation involves reaction of an unprotected peptide segment bearing a C-terminal α-thioester with a second unprotected peptide segment containing an N-terminal Cysteine residue. Thiol
35 exchange yields a thioester-linked intermediate which spontaneously rearranges to a native

amide bond at the ligation site. We have determined that a peptide segment bearing an N-terminal Cysteine and a C-terminal thioacid is sufficiently stable under native ligation conditions that it requires no temporary protection of the C-terminal thioacid functionality. Accordingly, these peptide segments can be used as the middle segments in a sequential 5 ligation scheme involving three or more peptide segments as shown in FIG. 1. Once such a middle segment has ligated to the solid phase-bound thioester-containing peptide to generate a solid phase-bound peptide thioacid, the thioacid is easily converted to a thioester and can be reacted with the N-terminal Cysteine of the next peptide segment to be ligated. Alternatively, the incoming peptide segment may have an internal amino acid with a nonnatural side chain 10 bearing amino and thiol moieties on adjacent α atoms, i.e. in a 1,2 relation to one another, and an unreactive, unprotected non-cysteine amino acid residue at its N-terminus, which would lead to a nonlinear assembled peptide. Multiple ligations of distinct peptide segments to form an assembled peptide bound to the solid phase are contemplated. Once all ligations are complete, the linker binding the solid phase-bound peptide to the solid phase is cleaved, 15 releasing the assembled peptide, i.e., the full length peptide. This technique is applied to the total chemical synthesis of a random peptide of artificial sequence (Table 1 in Examples Section), and human Macrophage Migration Inhibitory Factor (MIF), a 115 amino acid cytokine involved in immune system function. See FIG. 16-20.

20 A. Peptide Synthesis

Peptide segments were synthesized in stepwise fashion by established machine-assisted solid-phase methods on polystyrene resins using *in situ* neutralization/HBTU activation protocols for Boc chemistry (L. Canne et al., Tetrahedron Lett. 38:3361-3364 (1997)) on Boc-aminoacyl-OCH₂-PAM resins, thioester-generating resins (Hojo, et al., Bull. 25 Chem. Soc. Jpn. 64:111-117 (1991)), or thioacid-generating resins. After chain assembly was complete, peptides were deprotected and simultaneously cleaved from the resin by treatment with anhydrous HF containing 5% *p*-cresol, lyophilized, and purified by preparative HPLC. The N-terminal peptide segment was modified prior to HF cleavage as outlined in FIG. 17A.

30

B. Preparation of the Solid Phase

The solid phase is prepared as depicted in FIG. 5A and 5B. FIG. 5A is a scheme for preparing PEGA resin as a solid support. FIG. 5B is a generalized diagram for the preparation of any solid phase. An amino-Spherilose™ (Isco) affinity resin was derivatized 35 with Boc-aminoxyacetic acid as shown in FIG. 17B.

Other resins to be used as the solid phase include EAH Sepharose (Pharmacia), Amino PEGA (Novabiochem), CLEAR base resin (Peptides International), long chain alkylamine controlled pore glass (Sigma), HCl•PEG polystyrene (PerSeptive Biosystems), Lysine Hyper D resin (Biosepra), ArgoGel Base resin (Argonaut Technologies). These resins 5 are available in amino-derivatized form or are readily converted to amino-derivatized form.

C. Coupling of Modified N-terminal Peptide Segment to Solid Phase.

The modified peptide, containing a ketone moiety, as depicted in FIG. 17A, is 10 dissolved in 6M guanidine• HCl, 0.1M Na acetate, 0.15M methionine, pH 4.6 (1.6mM) and added to the aminoxy functionalized solid support, which had previously been thoroughly washed in the same buffer, and allowed to react at room temperature overnight (FIG. 16A, Step #1).

15 II. LIGATION IN THE N- TO C-TERMINAL DIRECTION

A. Ligation Reactions. The peptide segment to be ligated to the resin-bound peptide thioester was dissolved in 6M guanidine• HC1, 0.1M Na acetate, 0.15M methionine, 0.5% thiophenol, pH 7.5 (3.7-4.0mM) and added to the resin bound peptide thioester, which was thoroughly washed in the same buffer, and allowed to react at room temperature 20 overnight (FIG. 16A and 16B, Steps #2 and 4). Preferably the concentration of the first peptide segment can range from 1 to 150 mM; more preferably from 5-100 mM, most preferably from 10-50 mM, depending on the particular peptide segment.

One of skill in the art will understand that concentrations of the first peptide segment and the second and other incoming peptide segments can be optimized using routine 25 experimentation. Concentrations of the second and additional incoming peptide segments can range from 1-200 mM, more preferably from 5-100 mM, and most preferably from 10-59 mM, depending on the particular peptide segment.

Excess first peptide segment and/or excess incoming peptide segments can be readily removed from the solid phase bound peptide by filtration and washing.

30

B. Conversion of Thioacid to Thioester using Bromoacetic Acid or Iodoacetic Acid.

The use of Bromoacetic acid or Iodoacetic acid is an improved method of generating

peptide- α COSR thioesters from peptide- α COSH thioacids. In order to insure solubility of long unprotected peptides, 6 M guanidine-HCl at near pH 4 is used. Reactions is carried out near pH 4. Under such conditions, the only group reactive with Bromoacetic acid or Iodoacetic acid is the thioacid. Benzyl bromide, a hydrophobic compound, does not dissolve 5 completely in solution, resulting in slow and heterogeneous reactions. The advantages of using bromoacetic acid or iodoacetic acid are that both are readily soluble in 6 M guanidine-HCl (an aqueous solution) at near pH 4, both result in quick completion of the desired reaction, both elute in the void volume of typical reverse-phase HPLC, and allow processing of large amounts of peptide segments.

10 The resin-bound peptide thioacid is thoroughly washed in 6M guanidine• HCl, 0.1M Na acetate, 0.15M methionine, pH 4.6 and treated with a 50mM solution of bromoacetic acid in the same buffer for 15 min, followed by thorough washing with the pH 4.6 buffer (FIG. 16B, Step #3).

15 C. Cleavage from the Solid Phase.

Cleavable handles useful in the ligations in the N- to C-terminal direction must be capable of being stable to ligation conditions, stable to stepwise solid phase chemistries, able to be covalently linked in unprotected form to the solid phase, and be cleavable without damaging the assembled polypeptide. Any cleavable handles satisfying these requirements 20 can be used, including, but not limited to: MSC handle, photolabile linkers, CAM esters (-OCHCONH-), (-O-CH₂-Ø-SO-CH₂-CO-), (-O-CRH-CO-Ø-O-CH₂-CO-). For example, (-O-CH₂-Ø-SO-CH₂-CO-) may be used as a handle cleavable under any of the following conditions: (1) HF, DMS; (2) SciCl₄, TFA; or red of Sulfoxide and TFA cleavage; (3) NaOH, water; or (4) red of sulfoxide and TBAF in DMF. See Samamen, J. M., J. Org. 25 Chem. 53:561 (1988). As another example, the (-O-CRH-CO-Ø-O-CH₂-CO-) may be used as a cleavable handle under any of the following conditions: (1)NaOH, water(CAM Linker); (2) ZnCH₃COOH/Water; (3) photolysis. See Tjoeng et al., Synthesis 897 (1981); Sheehan et al., J. Org. Chem. 38:3771 (1973); Serebryakov et al., Tetrahedron 34:345 (1978); Hendrickson et al., Tetrahedron Lett. 343 (1970); Ceccato, M.L. et al., Tetrahedron Lett. 30 31:6189-6192 (1990); J. Martinez et al., Tetrahedron Lett. 41:739 (1985). One of skill in the

art will readily appreciate the suitability of known cleavable handles for the purposes described herein.

The following conditions can be used for cleavage of the linker to release the assembled polypeptide from the solid phase, particularly when an MSC handle is used.

5 Aliquots of resin-bound peptide are treated with 6M guanidine• HCl, 0.1M Na acetate, 0.15M methionine, containing 200 mM hydrazine, at pH ~14 for 2 min, followed by washing with an equal amount of 6M guanidine• HCl, 0.1M Na acetate, 0.15M methionine, pH ~2 and an equal amount of 6M guanidine• HCl, 0.1M Na acetate, 0.15M methionine, pH 4.6. The combined eluants of free peptide are analyzed by analytical HPLC and electrospray mass

10 spectrometry (FIG. 16B, Step #5).

II. SOLID PHASE LIGATIONS IN THE C- TO N-TERMINAL DIRECTION.

The discussion regarding N- to C-terminal ligations above applies equally well to C- to N-terminal ligations, except, as shown in FIG. 23, that: (1) the first peptide segment is bound to the solid phase via its C-terminus, i.e. the C-terminal peptide segment of the resulting assembled polypeptide is the one modified with a cleavable handle and (2) the incoming (i.e. second, third, additional) peptide segments do require temporary protection of their N-terminal Cysteine (see steps 2-4). Optionally, all Cysteine residues of the incoming or middle peptide segments can be temporarily protected along with the N-terminal Cysteine.

20 As outlined in the scheme (FIG. 23), the C-terminal peptide segment bearing a cleavable handle is coupled to the solid support by reaction with a corresponding functional group on the solid support (e.g. resin), for example, through an oxime linkage (aminoxyacetyl group on the resin and a ketone [via levulinic acid] on the peptide), or the reverse (aminoxyacetyl group on the peptide and a ketone on the solid phase).

25 Once the first peptide segment is bound to the solid phase as shown in step 1 of FIG. 21, the incoming (second) peptide segment, comprising an N-terminal protected Cys (PG-Cys) and a C-terminal thioester, reacts with the N-terminal unprotected Cys of the resin-bound first peptide segment through the native chemical ligation reaction. After ligation is complete, the protecting group of the N-terminal Cys is removed (step 3 of FIG. 21), and the

next peptide segment is added (step 4/2 of FIG. 21). Once all ligations are complete (step 5 of FIG. 21), the handle attaching the sequentially ligated peptide to the resin is cleaved, releasing the full length peptide. This C- to N-terminal technique is applied to the total chemical synthesis of a random peptide of artificial sequence and to human secretory 5 phospholipase A2, group 5 ("PLA2G5"), a 118 amino acid enzyme, as described below.

A. Peptide Synthesis

Peptide synthesis for solid phase sequential native chemical ligation in the C- to N-terminal direction is essentially the same as described above for solid phase sequential native 10 chemical ligation in the N- to C-terminal direction.

See Example 7 below for details re stepwise solid phase peptide synthesis of the peptide segments.

B. Preparation of the Solid Phase

15 Preparation of the solid phase for the C-to N-terminal direction is identical to that described for the N- to C-terminal direction.

C. Coupling of the Modified C-Terminal Peptide Segment to Solid Phase

Conditions for coupling the modified C-terminal peptide segment to the solid support 20 can be identical to that outlined for coupling of the modified N-terminal peptide in the N- to C-terminal ligations as described above.

D. Ligation in the C- to N-terminal Direction

Conditions for the native chemical ligation reactions in the C- to N-terminal direction 25 can be identical to that outlined for N- to C-terminal ligations as described above, except that

the N-terminal cysteine containing peptide segment is solid phase bound and the incoming thioester containing peptide segment is in solution.

E. Cysteine Protecting Groups and Removal

5 Any of the known protecting groups suitable for protecting the N-terminal Cys of a peptide segment can be used, provided that they are stable to ligation conditions, stable to conditions for adding the linker, and removable from the peptide segment under conditions that are not harmful to the solid-phase bound peptide, the linker, the resin, or the cleavable handle, if used. The protecting groups must also be stable to stepwise solid phase peptide synthesis conditions. An example of a protecting group is ACM (Acetamidomethyl), which provides cysteine side chain protection (-SCH₂NHCOCH₃), and can be cleaved with mercury(II)acetate, or other suitable reagents. Fmoc (9Fluorenylmethylcarbamate) provides alpha amino protection, can be cleaved in 20% piperidine in DMF and works well with hydrophilic peptides. DNPE (2-(2,4-dinitrophenyl)ethyl) provides cysteine side chain protection and cleaves in 50% piperidine in DMF. Para-nitrobenzensulfonyl provides alpha-amino protection, and is cleaved in 1 M DBU/1 M beta-mercaptoethanol in DMF. Additional cysteine protecting groups include, but are not limited to, Sulfmoc, NSC, Dde, Boc-Cys(Acm)-OH, Fmoc-Cys-(Mob)-OH, Boc-Cys(Fm)-OH, and Boc-Cys(DNPE)-OH, wherein Acm=acetamidomethyl, Mob = methoxybenzyl, Dnpe = 2-(2,4-dinitrophenyl)ethyl, Fm = 9-fluorenylmethyl. See Protective Groups inOrganic Synthesis, Green, T.W. and Wuts, P.G.M. eds, (2d Ed. 1991), particularly p. 293-294, 318-319; R. Merrifield, J. Org. Chem. 43:4808-4816 (1978); V.V. Samukov et al., Tetrahedron Lett. 35:7821-7824 (1994); B.W. Bycroft et al., J. Chem. Soc. Chem. Comm. 776-777 (1993); M. Royo et al., Tetrahedron Lett., 33:2391-2394 (1992); S.C. Miller, J. Am. Chem. Soc. 119:2301-2302 (1997). Certain 25 protecting groups can make peptide segments insoluble. For example, certain hydrophobic peptide segments may become insoluble upon addition of a protecting group. One of

ordinary skill in the art can readily ascertain the suitability of any particular protecting group for a peptide segment.

Removal of Fmoc as a Cys Protecting Group. One embodiment involves removal 5 of an Fmoc protecting group from the N-terminal Cys of a solid-phase bound peptide. After ligation with a peptide with an N-terminal Fmoc-Cys, the resin bound peptide is washed with 6 M guanidine•HCl, 0.1 M NaPi, 0.15 M methionine, pH 7, followed by water, followed by DMF. The resin is then treated with two aliquots of 20% piperidine in DMF, 5 minutes each. The resin is then washed thoroughly with DMF, followed by water, followed by 6 M 10 guanidine•HCl, 0.1 M NaPi, 0.15 M methionine, pH 7.

Removal of ACM as a Cys Protecting Group. After ligation with a peptide with an N-terminal Cys(ACM), the resin bound peptide is washed with 6 M guanidine•HCl, 0.1 M NaPi, 0.15 M methionine, pH 7, followed by 3% aqueous acetic acid. The resin is then treated with a solution of mercury(II)acetate in 3% aqueous acetic acid (15mgs/ml) for 30 15 minutes, followed by washing with 3% aqueous acetic acid. The resin is then washed with 6 M guanidine•HCl, 0.1 M NaPi, 0.15 M methionine, pH 7, followed by treatment with 20% beta-mercaptoethanol in 6 M guanidine•HCl, 0.1 M NaPi, 0.15 M methionine, pH 7 for 30 min. The resin is then washed with 6 M guanidine•HCl, 0.1 M NaPi, 0.15 M methionine, pH 7.

20

F. Cleavage from the Solid Phase

Cleavable handles are used to cleave the solid-phase bound peptide from the solid phase for ligations in the N- to C-terminal direction, in the C- to N-terminal direction, and in the bidirectional approach (both N- to C-terminal ligation and C- to N-terminal ligation). For 25 solid phase sequential native chemical ligations in the C- to N-terminal direction (and for bidirectional ligations using C- to N-terminal ligation), the requirements of cleavable handle

are the same as for those useful in the N- to C-terminal direction, with the additional requirement that the cleavable handle be stable under conditions used for removal of the protecting group from the N-terminal cysteine of the solid-phase bound peptide.

Cleavage of a peptide-CAM ester linkage to the solid phase. Aliquots of resin-bound peptide are washed with 8M urea, 0.1M NaPi, pH 7, followed by treatment for 2 minutes with 0.25N NaOH in the same 8M urea buffer (resulting pH~14). The resin is then washed with an equal amount of 0.25N HCl in the same 8M urea buffer (resulting pH~2), followed by thorough washing with the 8M urea buffer. The combined eluants of free peptide are analyzed by HPLC and electrospray mass spectrometry.

10

III. BIDIRECTIONAL SOLID PHASE SEQUENTIAL NATIVE CHEMICAL LIGATION.

Yet another embodiment of the invention relates to bidirectional solid phase protein synthesis that incorporates aspects of both the N- to C-terminus and C- to N-terminus sequential solid phase protein synthesis approaches. In the bidirectional approach, a peptide segment having either or both an N-terminal Cysteine and/or a C-terminal thioester is attached to a solid phase via a side chain of one of its amino acid residues. See FIG. 25A, B, C. The peptide segment can then be ligated at either terminus to a second peptide segment, followed by ligation at the other terminus to a third peptide segment. In this bidirectional approach, if the peptide segment attached to the solid phase has both a protected N-terminal Cysteine and a C-terminal thioester, second and third peptide segments can be added at both ends in subsequent ligations. Additional peptide segments can then be added at either end of the ligated, solid phase bound peptide. The ligations in either direction are accomplished using the methods described herein for ligations in either the C- to N-terminal direction or the N- to C-terminal direction.

25 Alternatively, the first peptide segment attached via one of its internal amino acid residues to the solid phase can be used for only uni-directional ligations. For example, the

peptide segment attached to the solid phase can be ligated to a second peptide segment at one terminus, followed by one or more ligations to additional peptide segments at the same terminus of the second peptide segment. In this embodiment, the peptide segment bound to the solid phase can be used for either sequential solid phase native chemical ligations in the

5 C- to N-terminal direction or for sequential solid phase native chemical ligations in the N- to C-terminal direction. In this embodiment, the peptide segment bound to the solid phase can be bidirectionally capable (i.e. having both a protected N-terminal Cysteine and a C-terminal thioester) while being used for unidirectional sequential ligations (i.e. having either a protected N-terminal Cysteine or a C-terminal thioester).

10 The first peptide segment is bound to the solid phase via a side chain of one of its amino acid residues, which is bound to a cleavable handle, which is bound to the solid phase via a functional chemical moiety that is capable of chemoselectively forming a covalent bond with a complementary functional chemical moiety on the solid phase, as illustrated in FIG. 25.

15 For example, the first peptide segment can be bound to the solid phase via the side chains of a lysine, aspartic acid or glutamic acid, in which case a cleavable handle based on functionalities, such as allyloxycarbonyl (alloc) or Fmoc, i.e. cleavable under orthogonal conditions, may be used to connect the peptide segment to the solid phase via the side chain of its lysine, aspartic acid or glutamic acid. As another example, an oxime bond may be 20 formed by the first peptide segment and the solid phase, wherein the first peptide segment comprises either an amino-oxy or ketone chemoselective functional group and the solid phase comprises a complementary chemoselective functional group, such as a ketone or amino-oxy, respectively.

25 IV. Use of Cleavable Linkers and Mass Spectrometry to Monitor Ligation Reactions

Various known cleavable linkers can be used to monitor the solid phase sequential ligations. These cleavable linkers are placed between the solid phase and the first peptide segment which is covalently bound to the cleavable handle, e.g. solid phase — cleavable linker—cleavable handle—peptide segment. The cleavable linkers are capable of being 30 readily cleaved to permit mass spectrometric analysis of a small portion of solid phase-bound peptide to monitor the coupling and ligation reactions.

For example, when the solid phase consists of resin beads, one can take a few resin beads from the reaction mixture after the coupling reaction or after each ligation reaction to determine the extent of reaction. Particularly preferred cleavable linkers include photolabile cleavable linkers for MALDI mass spectrometry, including 3-nitro-4(methylamino)benzoyl-.

- 5 See FIG. 5A. A small aliquot of the reaction mixture is removed for MALDI MS analysis and dried on a slide in mixture with a matrix solution. The laser of the MALDI mass spectrometer cleaves the photolabile linker on the mass spectrometer's stage, permitting mass analysis of the released peptides.

Another preferred cleavable linker is one that is cleavable by TFA (trifluoroacetic acid), which is useful for electrospray ionization mass spectrometry. With TFA-cleavable 10 linkers, the peptides are cleaved from the solid phase prior to ESI MS.

EXAMPLES

Example 1: Preparation of the Solid Phase for N- to C-Terminal Ligations

The preparation of the solid phase is schematically diagrammed in FIG. 5. The solid phase is a resin, for example, Amino PEGA (0.2-0.4 mmol/g swelled in methanol) or an amino-Spherilose affinity resin (15-20 Tmol/ml, 0.6-0.9 mmol/g dry resin), available from Pharmacia, NovaSyn or Isco. The resin (PEGA or Isco) is washed with DMF (dimethylformamide), then is washed briefly with 10% DIEA (diisopropyl ethylamine). Two 30 sec DMF flow washes are used. A photocleavable linker (PCL) (See FIG. 5A) is activated with one equivalent of HBTU (2-(1H-benzotriazol-1-yl)-1,1,3,3-tetramethyl-uronium hexafluorophosphate) and DIEA in DMF for 5-10 min). This activated photocleavable linker is then added to the resin and is left standing at room temperature for ~3 hrs (ninhydrin can be used with Isco).

Two 30 sec. DMF flow washes are used, followed by TFA (1 min x 2), and two more 30 sec. DMF flow washes. The remaining steps are in abbreviated form:

- 15 - 10% DIEA (1 min x 2)
 - DMF flow wash (30 sec x 2)
 - addition of activated Boc-aminoxyacetic acid (activated with one equivalent DIC and N-hydroxysuccinimide in DMF for 30-60 min)
 - left standing at room temperature for ~1 hr (ninhydrin can be used with Isco)
- 20 - DMF flow wash (30 sec x 2) [resin can be stored at this stage]
 - TFA (1 min x 30)
 - DMF flow wash (30 sec x 2)
 - 10% DIEA (1 in x 2)
 - DMF flow wash (30 sec x 2)
- 25 - thorough washing with aqueous buffer (6 M GuHCl, 0.1 M Na Acetate, pH 4.6) (1 ml x 5)

Example 2: Preparation of the First Unprotected Peptide Segment for N- to C-terminal Ligations

30 The following procedures are used to prepare the first peptide segment (N-terminus), which is diagrammed in FIG. 6, 7A and 7B.

- The peptide-resin is swelled in DMF
- TFA (1 min x 2)
- DMF flow wash (30 sec x 2)
- 35 - 10% DIEA (1 min x 2)

- DMF flow wash (30 sec x 2)
- Addition of MSC handle in DMF
- leave standing at room temperature for 1 hr
- add DIEA and leave standing for another hr
- 5 - use ninhydrin test to verify adequate coupling
- DMF flow wash (30 sec x 2)
- TFA (1 min x 2)
- DMF flow wash (30 sec x 2)
- 10% DIEA (1 min x 2)
- 10 - DMF flow wash (30 sec x 2)
 - addition of activated levulinic acid (activated as the symmetric anhydride with 0.5 equivalents of DIC in DCM for 5-10 min)
 - leave standing at room temperature for 30 min
 - ninhydrin test to verify adequate ligating
- 15 - DMF flow wash (30 sec x 2)
 - thorough washing with DCM
 - dry on lyophilizer
 - HF cleavage at 0° C for 1 hr using *p*-cresol as a scavenger
 - trituration and washing with cold ethyl acetate
- 20 - dissolve in 50% B and lyophilize
 - purify by preparative HPLC

Table 1

Solid Phase Sequential Ligations: N- to C-Terminal

3-Random Peptide Segment Model System

Lev-MSC-LTEGLHGFHVHEFGDNTAGCTSAGPHFNPLSRKHG-COS (1)

+ Resin-*PCL-ONH*₂

↓ 1. pH 4.6, 6M GuHCl, 0.1 M acetate

Resin-*PCL-oxime-MSC-LTEGLHGFHVHEFGDNTAGCTSAGPHFNPLSRKHG-COS* (1)

↓ 2. pH 4.6, 6M GuHCl, 0.1 M acetate, 50 mM BrAcOH

Resin-*PCL-oxime-MSC-LTEGLHGFHVHEFGDNTAGCTSAGPHFNPLSRKHG-COSAc* (1)

+ *H-CGFRVREFGDNTA-COS* (2)

↓ 3. pH 7.5, 6M GuHCl, 0.1M phosphate, 0.5% thiophenol

Resin-*PCL-oxime-MSC-LTEGLHGFHVHEFGDNTAGCTSAGPHFNPLSRKHG CGFRVREF-GDNTA-COS* (1+2)

↓ 4. pH 4.6, 6M GuHCl, 0.1M acetate, 50mM BrAcOH

Resin-*PCL-oxime-MSC-LTEGLHGFHVHEFGDNTAGCTSAGPHFNPLSRKHG CGFRVREF-GDNTA-COSAc* (1+2)

+ *H-CADPSEEVQKYVSDLELSA-OH* (3)

↓ 5. pH 7.5, 6M GuHCl, 0.1M phosphate, 0.5% thiophenol

Resin-*PCL-oxime-MSC-LTEGLHGFHVHEFGDNTAGCTSAGPHFNPLSRKHG CGFRVREF-GDNTACADPSEEVQKYVSDLELSA-OH* (1+2+3)

↓ 6. pH 14, 6M GuHCl, 0.1M phosphate, 200mM hydrazine

H-LTEGLHGFHVHEFGDNTAGCTSAGPHFNPLSRKHGCGFRVREF- GDNTACADPSEEVQKYVSDLELSA-OH (1+2+3)

PCL= photocleavable linker

Example 3: Solid Phase Native Chemical Ligation of Random Peptide Segments in Aqueous Solution in the N- to C-terminus direction.

The following procedures are used for solid phase ligations in the N- to C-terminus direction, as diagrammed in Table 1. General principals of native chemical ligation are 5 described in WO 96/34878, PCT/US95/05668, incorporated herein by reference.

The resin is washed with 6 M guanidine•HCl, 0.1 M Na Acetate, pH 4.6 (1 ml x 5) and drained. The modified N-terminal peptide segment is dissolved in 6 M guanidine•HCl, 0.1 M Na Acetate, pH 4.6 and added to resin and is left standing at room temperature overnight. (The concentration of the first segment is at least 5 mM). The next morning, 10 resin is washed with 6 M guanidine•HCl, 0.1 M Na Acetate, pH 4.6 (1 ml x 5) and drained. A sample of resin is removed for MALDI MS analysis and is washed with 50% B, MeOH, DCM and dried. A sample of resin is removed for base cleavage and is treated with 200 Φ l 6 M guanidine•HCl, 0.1 M Na Pi, 200mM hydrazine, pH ~14 for 2 min and drained, resin is washed with 200 Φ l 6 M guanidine•HCl, 0.1 M Na acetate, 200mM hydrazine, pH ~2 and 15 with 200 Φ l 6 M guanidine•HCl, 0.1 M Na Acetate, pH 4.6 and the combined eluants treated with TCEP prior to injection on HPLC.

In preparation for addition of the next peptide segment, the resin is washed with 6 M guanidine•HCl, 0.1 M Na Pi, pH 7.5 (1 ml x 5) and drained. The second peptide segment (Cys--COSH) is dissolved in 6 M guanidine•HCl, 0.1 M Na Pi, pH 7.5, 0.5% thiophenol and 20 added to resin. This mixture is left standing at room temperature overnight. The next morning, the resin is washed with 6 M guanidine•HCl, 0.1 M Na Acetate, pH 4.6 (1 ml x 5) and drained. Samples of resin are removed for Maldi and base cleavage and treated as above

The solid phase-bound peptide is then converted from COSH to COSAc by treating the resin with 50 mM BrAcOH in 6 M guanidine•HCl, 0.1 M Na Acetate, pH 4.6 for 15 min.

25 The resin is washed with 6 M guanidine•HCl, 0.1 M Na Acetate, pH 4.6 (1 ml x 5) and drained.

In preparation for addition of the next peptide segment, the resin is washed with 6 M guanidine•HCl, 0.1 M Na Pi, pH 7.5 (1 ml x 5) and drained. The final peptide segment is dissolved in 6 M guanidine•HCl, 0.1 M Na Pi, pH 7.5, 0.5% thiophenol and added to resin. 30 This reaction mixture is left standing at room temperature overnight. The next morning, the resin is washed with 6 M guanidine•HCl, 0.1 M Na Acetate, pH 4.6 (1 ml x 5) and drained. A sample of resin are removed for monitoring by MALDI MS analysis.

The assembled peptide is removed from the solid phase via base cleavage of the cleavable handle from the remaining resin as outlined above only on a larger scale followed by purification by HPLC or desalting on PD-10 column and lyophilization.

5 Example 4: Solid Phase Native Chemical Ligation of C5a(1-74) (74aa) in the N- to C-Terminal Direction.

This example describes solid phase sequential native chemical ligation in the N- to C-terminal direction of C5a, Complement Factor 5A. The sequence of C5a is: (SEQ ID NO. ?)

TLQKKIEEIAAKYKJSVVKKCCYDGACVNNDTCEQRAARISLGPKCIKAFTECCVV

10 ASQLRANISHKDMQLGR.

This peptide is prepared using solid phase sequential native ligation of 3 peptide segments: C5a(1-20), C5a(21-46), and C5a(47-74). The procedures used to synthesize C5a by solid phase ligations are identical to those described in the solid phase sequential native ligation of MIF (See **Example 5**).

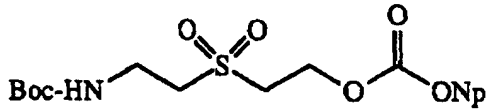
15

Example 5: Solid Phase Sequential Native Chemical Ligation of MIF(1-115) (115 aa) in the N-Terminal to C-Terminal Direction.

The sequence of MIF(1-115) is (SEQ.ID.NO.):
 MPMFIVNTNVPRASVPDGFLSELTQQLAQATGKPPQYIAVHVPDQLMAFGGSSEPC
 20 ALCSLHSIGKIGGAQNRSYSKLLCGLLAERLRISPDRVYINYYDMNAASVGWNNSTF
 A. This peptide is prepared using solid phase sequential native ligation of 3 peptide segments: MIF(1-59), MIF(60-80) and MIF(81-115). See **FIG. 16-20**.

Step #1: The first unprotected peptide segment, MIF(1-59) is coupled to a solid phase as depicted in **FIG. 18**. The coupling conditions are 6M guanidine• HCl, 0.1M
 25 NaAcetate, 0.15M Methionine, pH 4.6, 24 hours.

The MSC handle used is:



30

This cleavable handle is based on methylsulfonylethyoxy carbonyl (MSC) amine protecting group. It is easily added to unprotected amino terminus of peptide-resins, survives HF deprotection and cleavage from the resin, is quickly and cleanly cleaved by aqueous base, and is designed with a protected amine which can be derivatized with a variety of functionalities.

5 **Step #2:** The second unprotected peptide segment (Cys60-MIF(61-80)-COSH) is then ligated to the solid phase-bound first unprotected peptide segment, under the conditions 6 M guanidine• HCl, 0.1M NaPi, 0.5% thiophenol, 0.15M Methionine, pH 7.5, 24 hours.

Step #3: The solid phase-bound peptide, MIF(1-80)-COSH, is then activated to the thioester under the following conditions: 50 mM BrCH₂COOH, 6M guanidine• HCl, 0.1M 10 NaAcetate, 0.15M Methionine, pH 4.6, 15 min.

Step #4: The third unprotected peptide segment (Cys81-MIF82-115-COOH) is ligated to the solid phase-bound peptide with 6 M guanidine• HCl, 0.1M NaPi, 0.5% thiophenol, 0.15M Methionine, pH 7.5, 24 hours.

Step #5: The MIF(1-115) bound to the solid phase is then cleaved from the solid 15 support by base cleavage of the cleavable handle under the cleaving conditions: 6 M guanidine• HCl, 0.1M NaAcetate, 0.15M Methionine, 200 mM hydrazine, at pH-14 for 2 min., followed by 6 M guanidine• HCl, 0.1M NaAcetate, 0.15M Methionine, 200 mM hydrazine, at pH-2. The expected mass of the assembled peptide MIF(1-115) released upon 20 base cleavage is 12450 Da. FIG. 20C and 20D are mass spectra of the assembled peptide having an expected mass of 12450. FIG 20D is a reconstruction of the mass spectrum of FIG 20C. FIG 20B is an HPLC chromatogram of the assembled peptide.

Example 6: Solid Phase Native Chemical Ligation of Phospholipase A2, group 5(1-118) (118aa) in the C- to N-terminal Direction.

25 The sequence of Phospholipase A2, group 5 (PLA2G5) is: (SEQ ID NO:):

GLLDLKSMEKVTGKNALTNYGFYGCYCGWGRGTPKDGTDWCCWAHDHCYGRL
EEKGCNIRTQSYKYRFAWGVVTCCEPGPFCHVNLCACDRKLVYCLKRNLRSPNPQYQ
YFPNILCS.

This peptide is prepared using solid phase sequential native ligation of 4 peptide segments: 30 PLA2G5 (1-25), PLA2G5 (26-58), PLA2G5(59-87) and PLA2G5 (88-118). The procedures used to synthesize PLA2G5 by solid phase ligations are identical to those used for synthesizing the random sequence using ACM protection of the N-terminal Cys residues of the middle segments, as described in Example 9. See FIG. 22 for the reaction scheme. The

Cam ester derivative is synthesized and incorporated into the C-terminal peptide segment according to the diagrams in FIG. 23, 24/FIG. 27

Example 7: Preparation of Modified C-terminal Peptide Segment (on-resin CAM 5 linker synthesis) (FIG. 27)

- The commerical resin of choice (MBHA, any Boc-AA-OCH₂-Pam resin) is swelled in DMF
- TFA (1 min x 2) (not necessary if working with MBHA resin)
 - DMF flow wash (30 sec x 2)
 - addition of activated Boc-Lys(Fmoc)-OH (HBTU/DIEA activation), check for completion of
 - 10 reaction after 10-15 minutes by ninhydrin test
 - DMF flow wash (30 sec x 2)
 - TFA (1 min x 2)
 - DMF flow wash (30 sec x 2)
 - 10% DIEA in DMF (1 min x 2)
 - 15 -addition of activated bromoacetic acid (activated as the symmetric anhydride with 0.5 equivalents of DIC in DCM for 5-10 minutes), check for completion of reaction after 30 minutes
 - by ninhydrin test
 - DMF flow wash (30 sec x 2)
 - 20 -addition of first Boc-protected amino acid of the sequence(Boc-AA-OH) 2M in 20% DIEA in DMF. Leave standing at room temperature for 3 hrs.
 - DMF flow wash (30 sec x 2)
 - synthesize rest of the sequence by standard protocols for Boc chemistry
 - remove Fmoc group by treating with 20% piperidine in DMF (5 min x 2)
 - 25 -DMF flow wash (30 sec x 2)
 - addition of activated levulinic acid (activated as the symmetric anhydride with 0.5 equivalents of DIC in DCM for 5-10 min), check for completion of reaction after 30 minutes
 - by ninhydrin test

- DMF flow wash (30 sec x 2)
 - thorough washing with DCM
 - thoroughly dry resin
 - HF cleavage at 0°C for 1 hr using p-cresol as a scavenger
- 5 -trituration and washing with cold ethyl ether
- dissolve in aqueous HPLC buffer and lyophilize
 - purify by preparative HPLC

**Example 8: Solid Phase Native Chemical Ligation of Random Peptide Segments in the
10 C- to N-terminal Direction using Fmoc protection (See FIG. 28)**

The following procedures can be used for solid phase ligations in the C- to N-terminal direction, as diagramed in Table 2. By example, a random peptide of (SEQ ID NO: ?): ALTKYGFYGCYGRLEEKGCADRKNILA can be ligated in three peptide segments (from C- to N-terminal direction): segment 1= CADRKNILA; segment 2 = CYGRLEEKG; and

15 segment 3 = ALTKYGFYG.

The resin is washed with 6M Gu•HCL, 0.1M Na Acetate, pH 4.6 (1 ml x 5) and drained. The modified C-terminal peptide segment (first peptide segment) is dissolved in 6M Gu•HCL, 0.1M Na Acetate, pH 4.6 (5 mM first peptide segment) and added to the resin and is left standing at room temperature overnight. The resin is washed with 6M Gu•HCL, 0.1M

20 Na Acetate, pH 4.6 (1 ml x 5) and drained. A sample is removed for base cleavage and is treated with 8M urea, 0.1M NaPi, pH 7, treated for 2 minutes with 0.25N NaOH in the same 8M urea buffer (resulting pH~14), washed with an equal amount of 0.25N HCl in the same 8M urea buffer (resulting pH~2), and the combined eluants treated with TCEP prior to injection on HPLC.

25 In preparation for addition of the next segment, the resin is washed with 6M Gu•HCl, 0.1M NaPi, pH 7.0 (1 ml x 5) and drained. The second peptide segment (Fmoc-Cys-peptide-COSR) is dissolved in 6M Gu•HCl, 0.1M NaPi, pH 7.0, 0.5% thiophenol (to at least 10 mM

to 50 mM second peptide segment) and added to the resin. the mixture is left standing at room temperature overnight. The resin is washed with 6M Gu•HCl, 0.1M NaPi, pH 7.0 (1 ml x 5), water (1 ml x 5), DMF (1 ml x 5), and the Fmoc protecting group removed by treating with two aliquots of 20% piperidine in DMF (5 min each). The resin is then washed with 5 DMF (1 ml x 5), water (1 ml x 5), and 6M Gu•HCl, 0.1M NaPi, pH 7.0 (1 ml x 5). A sample of resin is removed and base cleaved as above.

The final peptide segment is dissolved in 6M Gu•HCl, 0.1M NaPi, pH 7.0, 0.5% thiophenol and added to the resin. This mixture is left standing at room temperature overnight. The resin is then washed with 6M Gu•HCl, 0.1M NaPi, pH 7.0 and the assembled 10 peptide is removed from the solid phase via base cleavage of the cleavable handle from the remaining resin as outlined above only on a larger scale followed by purification by HPLC or desalting on PD-10 column and lyophilization.

These methods can be applied to make any peptides having cysteine residues.

15 Example 8A: Solid Phase Native Chemical Ligation of Random Peptide Segments in the C- to N-terminal Direction using DNPE protection

DNPE (2-(2,4-dinitrophenylethyl)) is another cysteine side chain protecting group which can be used for ligations in the C- to N-terminal direction. Example 8 was repeated using DNPE as the protecting group. The conditions for solid phase chemical ligation of 20 random peptide segments in the C- to N-terminal direction were identical to those used for Example 8 above except that in the removal of the DNPE protecting group, 50% piperidine is used.

Example 9: Solid Phase Native Chemical Ligation of Random Peptide Segments in the 25 C- to N-terminal Direction using ACM protection

The following procedures are used for solid phase ligations in the C- to N-terminal direction, as diagramed in Table 3. The same random polypeptide described in the Example above is ligated.

The resin is washed with 6M Gu•HCl, 0.1M Na Acetate, pH 4.6 (1 ml x 5) and drained.

- 5 The modified C-terminal peptide segment is dissolved in 6M Gu•HCl, 0.1M Na Acetate, pH 4.6 and added to the resin and is left standing at room temperature overnight. The resin is washed with 6M Gu•HCl, 0.1M Na Acetate, pH 4.6 (1 ml x 5) and drained. A sample is removed for base cleavage and is treated with 8M urea, 0.1M NaPi, pH 7, treated for 2 minutes with 0.25N NaOH in the same 8M urea buffer (resulting pH~14), washed with an
10 equal amount of 0.25N HCl in the same 8M urea buffer (resulting pH~2), and the combined eluants treated with TCEP prior to injection on HPLC

In preparation for addition of the next segment, the resin is washed with 6M Gu•HCl, 0.1M NaPi, pH 7.0 (1 ml x 5) and drained. The second peptide segment (Fmoc-Cys-peptide-COSR) is dissolved in 6M Gu•HCl, 0.1M NaPi, pH 7.0, 0.5% thiophenol (to at least 10 mM
15 second peptide segment) and added to the resin. The mixture is left standing at room temperature overnight. The resin is washed with 6M Gu•HCl, 0.1M NaPi, pH 7.0 (1 ml x 5), 3% acetic acid in water (1 ml x 5), and the ACM protecting group removed by treating with mercury(II)acetate in 3% acetic acid in water (15 mgs/ml) for 30 min. The resin is then washed with 3% acetic acid in water (1 ml x 5), 6M Gu•HCl, 0.1M NaPi, pH 7.0 (1 ml x 5),
20 and treated with 20% beta-mercaptopethanol in 6M Gu•HCl, 0.1M NaPi, pH 7.0 for 30 min, followed by washing with 6M Gu•HCl, 0.1M NaPi, pH 7.0 (1 ml x 5). A sample of resin is removed and base cleaved as above.

The final peptide segment is dissolved in 6M Gu•HCl, 0.1M NaPi, pH 7.0, 0.5% thiophenol and added to the resin. This mixture is left standing at room temperature
25 overnight. The resin is then washed with 6M Gu•HCl, 0.1M NaPi, pH 7.0 and the assembled peptide is removed from the solid phase via base cleavage of the cleavable handle from the

remaining resin as outlined above only on a larger scale followed by purification by HPLC or desalting on PD-10 column and lyophilization.

Table 2

5		Polymer-Supported Ligations C- to N- Terminal Direction Fmoc Protection
10		<i>H-CADRKNILA-CAM-Lys(Levulinic acid)-NH₂</i> (1) + Resin- <i>ONH₂</i>
15		↓ 1. pH 4.6, 6M Gu-HCl, 0.1 acetate <i>H-CADRKNILA-CAM-Lys-oxime-Resin</i> (1) + <i>Fmoc-CYGRLEEKKG-COSR</i> (2)
20		↓ 2. pH 7.5, 6M Gu-HCl, 0.1M phosphate, 0.5% thiophenol <i>Fmoc-CYGRLEEKGCADRKNILA-CAM-Lys-oxime-Resin</i> (1+2) ↓ 3. 20% piperidine/DMF
25		<i>H-CYGRLEEKGCADRKNILA-CAM-Lys-oxime-Resin</i> (1+2) + <i>H-ALTKYGFYG-COSR</i> (3) ↓ 4. pH 7.5, 6M Gu-HCl, 0.1M phosphate, 0.5% thiophenol
30		<i>H- ALTKYGFYG CYGRLEEKGCADRKNILA-CAM-Lys-oxime-Resin</i> (1+2+3) ↓ 5. pH 14, 8M Urea, 0.1M phosphate, 0.25N NaOH <i>H- ALTKYGFYG CYGRLEEKGCADRKNILA-OH</i>
35		

Table 3

Polymer-Supported Ligations C- to N- Terminal Direction ACM Protection	
5	
10	<i>H-CADRKNILA-CAM-Lys(Levulinic acid)-NH₂</i> (1) + Resin- <i>ONH₂</i>
15	↓ 1. pH 4.6, 6M Gu-HCl, 0.1 acetate <i>H-CADRKNILA-CAM-Lys-oxime-Resin</i> (1) + <i>H-C(ACM)YGRLEEKKG-COSR</i> (2)
20	↓ 2. pH 7.5, 6M Gu-HCl, 0.1M phosphate, 0.5% thiophenol <i>H-C(ACM)YGRLEEKGCADRKNILA-CAM-Lys-oxime-Resin</i> (1+2)
25	↓ 3. a. mercury(II)acetate in 3% Aq. AcOH b. 20% mercaptoethanol in pH 7.5, 6M Gu-HCl, 0.1M phosphate <i>H-CYGRLEEKGCADRKNILA-CAM-Lys-oxime-Resin</i> (1+2)
30	+ <i>H-ALTKYGFYG-COSR</i> (3) ↓ 4. pH 7.5, 6M Gu-HCl, 0.1M phosphate, 0.5% thiophenol <i>H- ALTKYGFYGCYGRLEEKGCADRKNILA-CAM-Lys-oxime-Resin</i> (1+2+3)
35	↓ 5. pH 14, 8M Urea, 0.1M phosphate, 0.25N NaOH <i>H- ALTKYGFYGCYGRLEEKGCADRKNILA-OH</i>

Example 10: Bidirectional Solid Phase Sequential Native Chemical Ligation

This example illustrates one of the embodiments of the bidirectional solid phase protein ligation approach, namely the situation starting with a first peptide segment bound to the solid phase, wherein the first peptide segment is a "middle piece" of the target protein

desired, i.e. the first peptide segment, bound to the solid phase, is used for ligations at both its N-terminal Cysteine and its C-terminal thioester.

Starting with one of the middle pieces of the target protein, a cleavable linker is added to the side chain of one of the amino acid residues of the middle piece. The side chain of any 5 amino acid residue having a protectable functional group can be used, including, preferably Aspartic Acid or Glutamic Acid. Most preferably, a Lysine amino acid residue is used. For example, a CAM ester cleavable handle or any other carboxylic acid protecting group may be adapted to attach the first peptide segment to the solid phase through the side chain of Aspartic or Glutamic Acid. One of skill in the art will readily appreciate the necessary 10 chemistries for accomplishing this step.

For example, the synthesis of a first peptide segment to be attached to the solid phase via an internal amino acid is illustrated in FIG. 25C. Starting with an appropriate solid phase (thioester or thiacid generating), the first peptide segment is synthesized using standard Boc protocols until the Lysine residue of choice is reached. Using Boc chemistry, a Lysine with 15 its side chain amine protected with an Fmoc group (Boc-Lys(Fmoc)-OH) is inserted at the appropriate location during solid phase stepwise peptide synthesis, followed by continued synthesis to the end of the first peptide segment. The Fmoc protecting group is removed at the end of the stepwise peptide synthesis and the cleavable handle coupled to the side chain amine (step B of FIG. 25C).

20 This method is much the same as the procedure outlined in FIG. 24, with the following differences: the levulinic acid in step 4 is replaced with the cleavable handle and the 20% piperidine used to cleave the Fmoc group (also part of step 4) is replaced with a much smaller concentration of an alternative base such as 1,8-Diazabicyclo[5.4.0]undec-7-ene (DBU), e.g. 1-2 equivalents of DBU in DMF. The reason is the middle peptide 25 segments, regardless of whether they generate thioacids or thioesters upon cleavage from the

resin, are connected to the resin by a thioester which would be cleaved in the presence of 20% piperidine.

For this particular strategy, the MSC handle is preferred, although other cleavable handles can be used. Attachment to the side chain amine of a lysine residue and further 5 modification of the linker with an appropriate functional group capable of reacting with a corresponding group on the solid phase ligation resin would be generally as outlined in FIG. 17A, with the exception that the amine of the MSC handle should be protected with an Fmoc instead of a Boc group. Since attachment to the peptide segment is through an internal amino acid residue, the N-terminal amino acid would be Boc protected and it is not possible for the 10 N-terminal amino group and the amino group of the MSC cleavable handle to be protected by the same group. Removal of the Fmoc group on the MSC cleavable handle would also need to be done with DBU instead of piperidine. As in FIG. 17A, levulinic acid is preferred for coupling to the linker with a corresponding aminoxyacetyl group on the solid support (FIG. 17B).

15 Two versions of the first peptide segment to be coupled to the resin are described below.

First Version. The first peptide segment has an unprotected N-terminal cysteine and a C-terminal thioacid (FIG. 25A). The second peptide segment (step 2. in FIG. 25A), to be ligated to the first peptide segment, is a peptide with a C-terminal thioester and optionally a 20 protected N-terminal Cysteine (if additional C- to N-terminal ligations are desired), wherein the C-terminal thioester is capable of reacting with the N-terminal Cys of the first peptide segment (i.e. in the C- to N-terminal direction). This step can be multiply repeated with additional peptide segments added in the C- to N-terminal direction, if desired, provided that the internal incoming peptide segments each comprise a protected N-terminal Cysteine, 25 which can be deprotected according to the standard C- to N-terminal solid phase native chemical ligation steps outlined in FIG. 21 (the final peptide segment to be added at the N-

terminus of the resulting product need not have an N-terminal Cysteine). After ligation is complete, the C-terminal thioacid of the resulting solid-phase bound peptide (i.e. ligation product of first and second peptide segments) is then converted to a thioester with bromoacetic acid (as outlined in N- to C-terminal ligations in Table 1 and diagrammed as 5 step 3 of FIG 25A). The next step (step 4 of FIG.25A) comprises ligation of the solid-phase bound peptide to a third peptide segment with an N-terminal Cys. This step can optionally be repeated, to add additional incoming peptide segments in the N- to C-terminal direction, if desired, provided that the internal incoming peptide segments each comprise an unprotected N-terminal Cysteine and a C-terminal thioacid, with conversion of the thioacid to thioester 10 after the ligation is complete and prior to addition of the next peptide segment. The final peptide segment to be added at the C-terminus of the resulting product need not have a C-terminal thioacid.

One of skill in the art will appreciate that multiple ligations can subsequently be performed in both directions if the appropriate protecting groups and other appropriate 15 chemistries are used on the middle piece or the solid-phase bound peptide. These additional steps are identical to the strategies used for the individual directions, i.e. N-terminal unprotected Cys plus C-terminal thioester for the N- to C- direction and N-terminal Cys(ACM) plus C-terminal thioester for the C- to N-terminal direction. Assuming the MSC linker is used, cleavage of the full length product from the resin would be in basic solution 20 (pH 12-14) as outlined in step 6 in Table 1. However, the preferred approach is to complete all ligation steps necessary for one direction, followed by the ligation steps for the other direction. As long as the solid phase bound peptide has either a protected N-terminal Cysteine or a C-terminal thioacid, ligations can proceed in either direction provided that the appropriate strategies as described herein are followed. If the solid phase bound peptide has 25 both an unprotected N-terminal Cysteine and a C-terminal thioester, any attempts at ligating

to an additional incoming peptide segment will result in cyclization of the solid-phase bound peptide.

Second Version. The second version of this scheme involves starting with ligation in the N- to C-terminal direction, followed by ligation in the opposite direction, as shown in FIG. 25B. The first peptide segment to be coupled to the resin comprises a temporarily protected N-terminal Cys and a C-terminal thioester. The ligation of a second peptide segment to the first peptide segment is then in the N- to C-terminal direction. Any subsequent ligations in the C- to N-terminal direction would first require removal of the protecting group.

10 Except for the attachment of the first peptide segment to the solid support, this strategy merely combines the procedures for N- to C- and C- to N-terminal ligations (described above).

References

- 15 S. Funakoshi et al., Chemoselective one-step purification method for peptides synthesized by the solid-phase technique, Proc. Nat. Acad. Sci. USA, 88:6981-6985 (Aug. 1991).
- S. Funakoshi et al., Affinity purification method using a reversible biotinylation reagent for peptides synthesized by the solid-phase technique, J. Chromatog. 638:21-27
20 (1993).
- M. Mutter et al., Pseudo-prolines (ψ Pro) for accessing inaccessible peptides, Pept. Res. 8(3):145-153 (1995).
- M. Baca et al., Chemical ligation of cysteine-containing peptides: synthesis of a 22 kDa tethered dimer of HIV-1 protease, J. Am. Chem. Soc. 117(7): 1881-1887 (1995).
- 25 J. Camarero et al., Chemical Ligation of Unprotected Peptides Directly From a Solid Support, J. Peptide Res. 51: 303-316 (1998).
- L. Canne et al., Total Chemical Synthesis of a Unique Transcription Factor-Related Protein: cMyc-Max, J. Am. Chem. Soc. 117:2998-3007 (1995).
- C. Cho et al., An Unnatural Biopolymer, Science 261:1303-1305 (1993).

- P. Dawson et al., Synthesis of Proteins by Native Chemical Ligation, *Science* 266:776-779 (1994).
- N. Fotouhi et al., *J. Org. Chem.* 54:2803-2817 (1989).
- G. Barany and R.B. Merrifield, A New Amino Protecting Group Removal by Reduction. Chemistry of the Dithiasuccinoyl (Dts) Function, *J. Am. Chem. Soc.*, 99(22):7363-7365 (1977).
- C. Hennard and J. Tam, Sequential Orthogonal Coupling Strategy for the Synthesis of Biotin Tagged β Defensin, Abstract P118, Fifteenth American Peptide Symposium, June 14-19, 1997.
- 10 C. Hyde et al., Some difficult sequences made easy, A study of interchain association in solid-phase peptide synthesis, *Int. J. Peptide Protein Res.* 43:431-440 (1994).
- W. Lu et al., *Biochemistry*, 36(4):673-679 (1997).
- C.-F. Liu and J. Tam, Peptide segment ligation strategy without use of protecting groups, *Proc. Natl. Acad. Sci. USA*, 91: 6584-6588 (1994).
- 15 C.-F. Liu and J.Tam, Chemical ligation approach to form a peptide bond between unprotected peptide segments. Concept and model study, *J. Am.Chem. Soc.* 116(10):4149-4153 (1994). Schnolzer et al., *Science* 256:221-225 (1992)
- Rose et al. *J. Am Chem. Soc.* 116:30-34 (1994)
- Liu et al., *Proc. Natl. Acad. Sci. USA* 91:6584-6588 (1994).
- 20 Dawson et al. *Science* 266:77-779 (1994).
- PCT/US95/05668, WO 96/34878
- Sakakibara S., *Biopolymers (Peptide Science)*, 37:17-28 (1995).
- Tam et al., *PNAS USA*, 92:12485-12489 (1995).
- T. Muir, A Chemical approach to the construction of multimeric protein assemblies, *25 Structure* 3:649-652 (1995).
- R. Merrifield, Solid Phase Peptide Synthesis: The Synthesis of a Tetrapeptide, *J. Am. Chem. Soc.*, 85:2149-2154 (1963).
- H. Muramatsu et al., Localization of Heparin-Binding, Neurite Outgrowth and Antigenic Regions in Midkine Molecule, *Biochem. And Biophys. Res. Commn.* 203 30 (2):1131-1139 (1994).
- PCT/US94/07222, WO 95/00846, Published January 5, 1995.

All publications and patent applications mentioned in this specification are herein incorporated by reference to the same extent as if each individual publication or patent application was specifically and individually indicated to be incorporated by reference.

5 The invention now being fully described, it will be apparent to one of ordinary skill in the art that many changes and modifications can be made thereto without departing from the spirit or scope of the appended claims.

WHAT IS CLAIMED IS:

1. A method of producing an assembled peptide in aqueous solution and on a solid phase comprising:
 - a) binding an unprotected first peptide segment to a solid phase via a linker, wherein
5 said unprotected first peptide segment comprises an N-terminus and a thioester of the formula -COSR at its C-terminus, wherein said linker comprises a cleavable moiety and said unprotected first peptide segment is bound to said linker at said N-terminus;
 - b) ligating a second unprotected peptide segment to said first peptide segment bound
10 to said solid phase, wherein said second peptide segment comprises a cysteine at its N-terminus and a thioacid at its C-terminus, and wherein said N-terminal cysteine of said second peptide segment is capable of selectively ligating to said C-terminus of said solid phase-bound first peptide segment to form a solid phase-bound peptide comprising a thioacid at its C-terminus;
 - c) converting said C-terminal thioacid of said solid phase-bound peptide to an activated thioester of the formula -COSR,
15
 - d) repeating steps b) and c) with a third unprotected peptide segment; and
 - e) optionally repeating steps b) and c) with additional unprotected peptide segments.
20
2. The method of claim 1, further comprising, after step e):
 - f) cleaving said linker to release an assembled peptide.
25
3. The method of claim 1, wherein said assembled polypeptide is from 20 to 1000 amino acids in length.
4. The method of claim 1, wherein said solid support is a bead resin.
30 5. The method of claim 1, wherein said cleavable moiety is a cleavable handle.
6. The method of claim 1, wherein said cleavable moiety is a cleavable linker.
35 7. The method of claim 1, wherein said peptide segments range in size from 5 to 99 amino acid residues.

8. The method of claim 1, wherein said peptide segments are all prepared by stepwise solid phase synthesis.

5 9. The method of claim 1, wherein the last peptide segment to be ligated onto the solid phase-bound peptide is derived from recombinant DNA expression.

10. The method of claim 1, wherein at least one of said peptide segments comprises an unnatural backbone structure.

10

11. The method of claim 1, wherein said converting step is accomplished using bromoacetic acid.

12. The method of claim 1, further comprising:

15 f) monitoring the ligation reactions using mass spectrometric analysis.

13. A method of producing an assembled peptide comprising:

20 a) binding an unprotected first peptide segment to a solid phase via a linker, wherein said unprotected first peptide segment comprises an N-terminus and a thioacid of the formula -COSH at its C-terminus, wherein said linker comprises a cleavable moiety and said unprotected first peptide segment is bound to said linker at said N-terminus;

b) converting said C-terminal thioacid of said first peptide segment to a thioester of the formula -COSR;

25 c) ligating a second unprotected peptide segment to said first peptide segment bound to said solid phase, wherein said second peptide segment comprises a cysteine at its N-terminus and a thioacid at its C-terminus, and wherein said N-terminal cysteine of said second peptide segment is capable of selectively ligating to said C-terminus of said solid phase-bound first peptide segment to form a solid phase-bound peptide comprising a thioacid at its C-terminus;

30

d) converting said C-terminal thioacid of said solid phase-bound peptide to an activated thioester of the formula -COSR,

e) repeating steps c) and d) with a third unprotected peptide segment; and

35 f) optionally repeating steps c) and d) with additional unprotected peptide segments.

14. A method of preparing an assembled peptide comprising:

a) binding a first peptide segment to a solid phase via a linker, wherein said first peptide segment comprises an N-terminal Cysteine and a C-terminal residue capable of 5 binding to said linker, wherein said linker comprises a cleavable moiety and said first peptide segment is bound to said linker at said C-terminal residue;

b) ligating a second peptide segment to said first peptide segment bound to said solid phase, wherein said second peptide segment comprises a cysteine at its N-terminus and a thioester at its C-terminus, and wherein said N-terminal cysteine of said second peptide 10 segment is capable of binding to a protecting group; and wherein said C-terminal thioester of said second peptide segments binds to said N-terminal cysteine of solid phase-bound first peptide segment to form a solid phase-bound peptide comprising protected Cysteine at its N-terminus;

c) removing said protecting group;

15 d) repeating steps b) and c) at least once with another peptide segment;

e) ligating a final peptide segment to said solid phase-bound peptide, wherein said final peptide segment is unprotected and comprises a C-terminal thioester;

f) cleaving said cleavable moiety to release an assembled peptide from the solid phase.

20

15. The method of claim 14, further comprising: monitoring said ligation reactions at any step using mass spectrometric analysis of the solid phase-bound peptides.

16. A method of preparing an assembled peptide comprising:

25 a) binding a first peptide segment to a solid phase via a linker, wherein said first peptide segment comprises an N-terminal Cysteine and a C-terminal residue capable of binding to said linker, wherein said linker comprises a cleavable moiety and said first peptide segment is bound to said linker at said C-terminal residue;

30 b) ligating a second peptide segment to said first peptide segment bound to said solid phase, wherein said second peptide segment comprises a peptide comprising a C-terminal thioester, wherein said C-terminal thioester of said second peptide segment binds to said N-terminal cysteine of solid phase-bound first peptide segment to form a solid phase-bound peptide comprising protected Cysteine at its N-terminus.

17. A method of preparing an assembled peptide comprising:

a) binding a first peptide segment to a solid phase via a linker to form a solid phase-bound peptide,

5 a) ligating at least a second peptide segment to said solid phase-bound peptide in aqueous solution to form a solid phase-bound peptide.

18. The method of claim 17, wherein said solid phase is water-compatible.

19. The method of claim 17, wherein said aqueous solution comprises 1-8 M urea.

10 20. The method of claim 17, wherein said aqueous solution comprises 1-6 M guanidine-HCl.

21. The method of claim 17, wherein said aqueous solution comprises 10-60% acetonitrile in water.

22. The method of claim 17, wherein said aqueous solution comprises a mixed 15 aqueous/organic solvent.

23. A kit for preparing assembled polypeptides comprising:

a) a first unprotected peptide segment, comprising a thioester at its C-terminus and an N-terminus, wherein said first unprotected peptide segment is bound to a solid phase via a 20 linker comprising a cleavable moiety;

b) a set of second unprotected peptide segments, each comprising a thioacid at their C-termini and a cysteine at their N-termini, wherein each of said second unprotected peptide segments have the same number of amino acids; and

c) one or more sets of different unprotected peptide segments, each comprising a 25 thioester at their C-termini and a cysteine residue at their N-termini, wherein each of the members of each set have the same number of amino acids.

24. The kit of claim 12, wherein said set of second unprotected peptide segments is comprised of peptides having the same length, but different amino acid sequences.

30

25. The kit of claim 12, wherein said set of second unprotected peptides consists essentially of identical peptides.

26. The kit of claim 13, wherein said one or more sets of different unprotected peptides comprise at least one set of peptides having the same length but different amino acid sequences.

5 27. An apparatus for producing assembled polypeptides, comprising:

- a) a solid support, having bound thereto a first unprotected peptide having a thioester at its C-terminus and a cleavable linker at its N-terminus, wherein said unprotected peptide is bound to said solid support via a linker;
- 10 b) a set of second unprotected peptides, each comprising a thioacid at their C-termini and a cysteine residue at their N-termini; and
- c) one or more sets of different unprotected peptides, each comprising a thioacid at their C-termini and a cysteine residue at their N-termini.

15 28. A method of producing polypeptide libraries of solid phase sequentially ligated assembled peptides, comprising:

- a) covalently binding a set of unprotected first peptide segments to a solid support via a linker, wherein said linker comprises a cleavable moiety stable under ligating conditions and 20 said unprotected first peptide segments are each bound to said linker at its N-terminus and has a thioester at its C-terminus;
- b) introducing a set of second unprotected peptide segments, wherein each of said second segments each comprise a cysteine residue at its N-terminus and a thioacid at its C-terminus, 25 under conditions suitable to permit ligation between said first unprotected peptide segments and said second unprotected peptide segments to form a natively ligated polypeptide bound to said solid support, wherein said bound polypeptide comprises a thioacid at its C-terminus;
- c) converting said thioacid to a thioester;
- 30 d) optionally repeating steps b) and c) with additional sets of unprotected peptide segments.

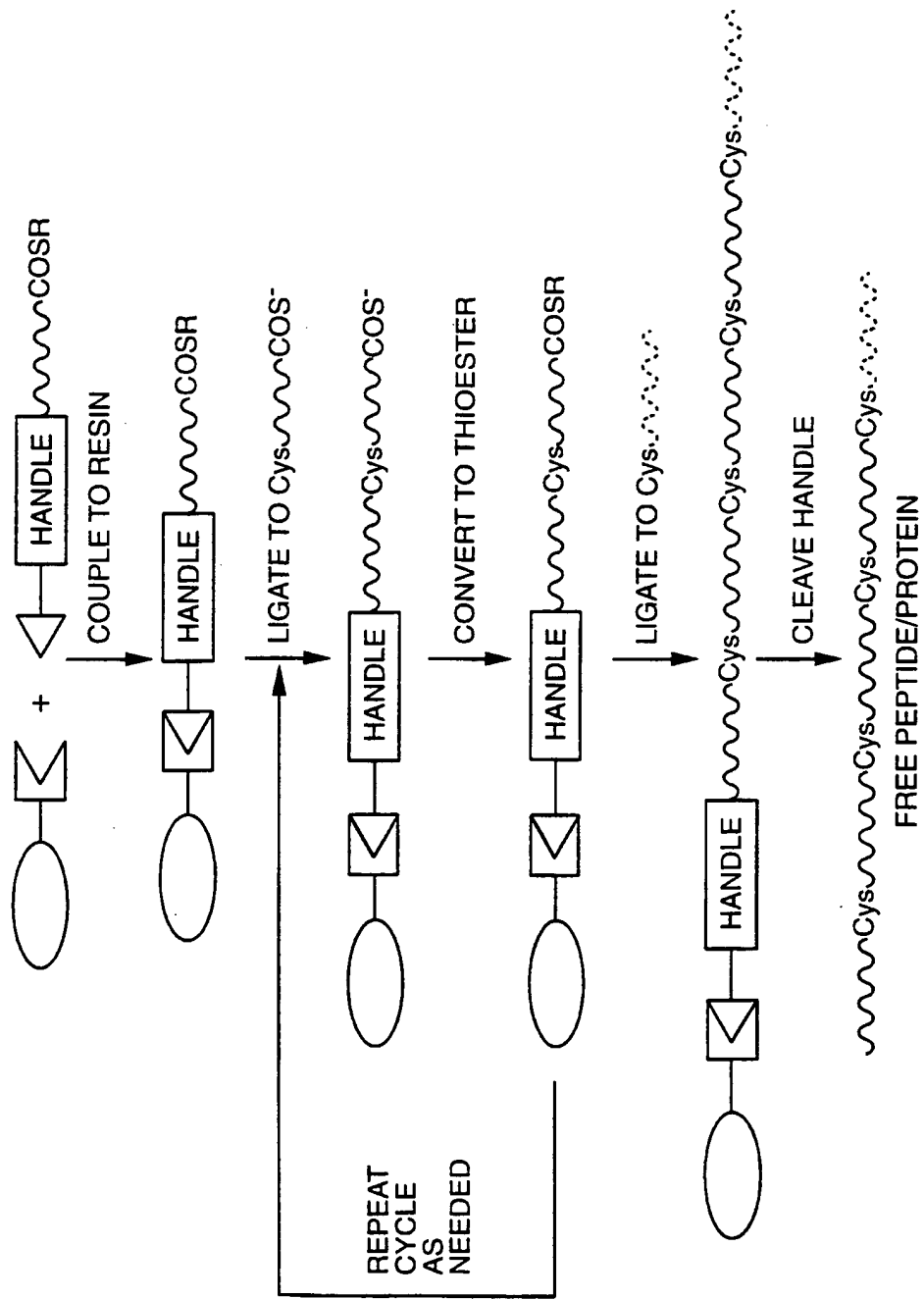


FIG. 1

2 / 31

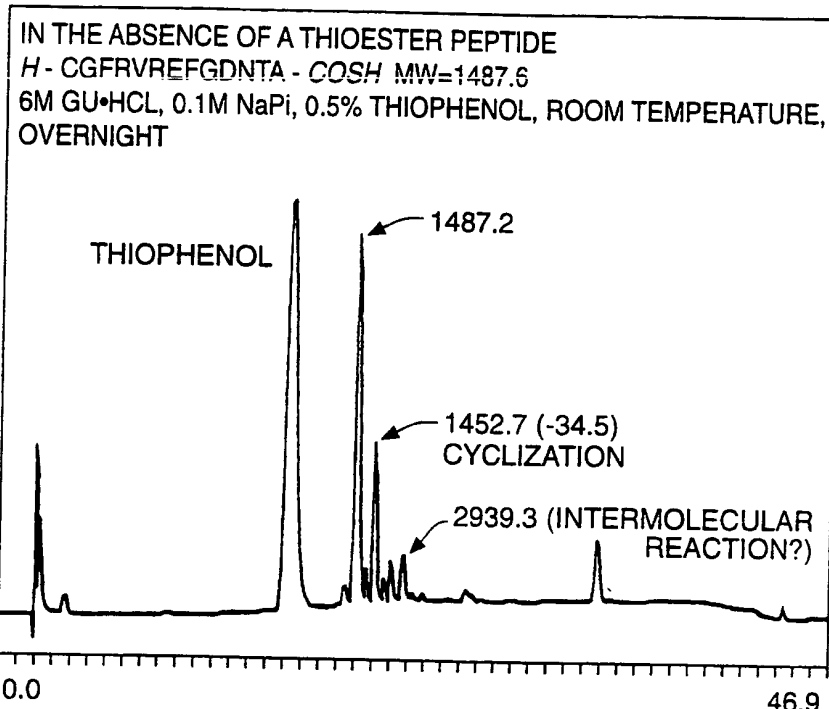


FIG._2A

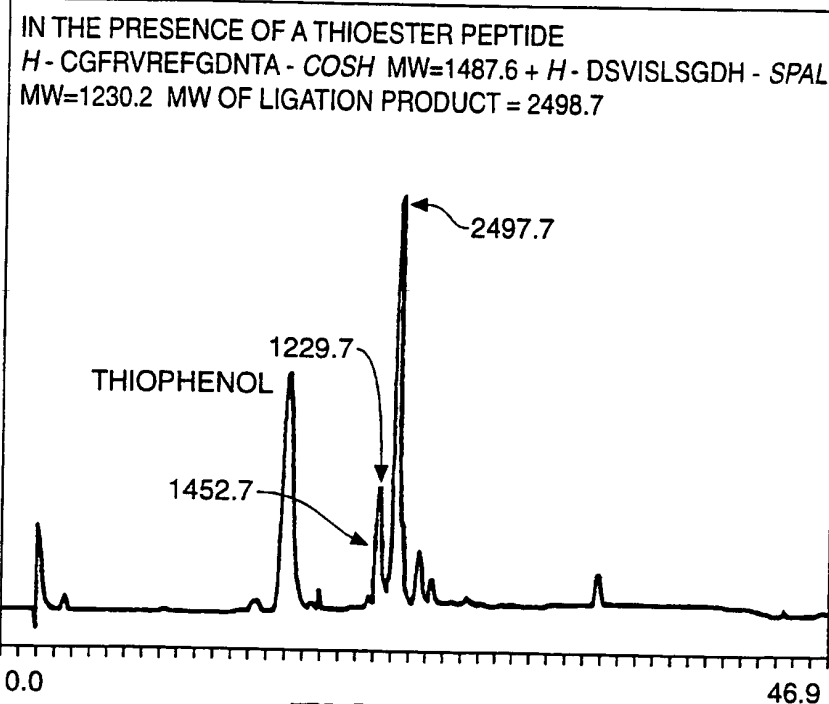


FIG._2B

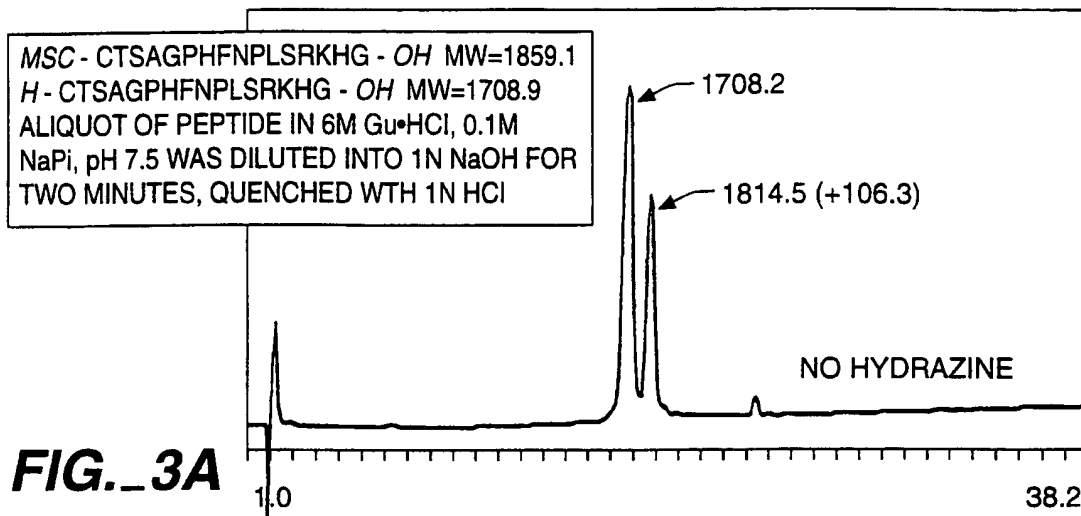


FIG._3A

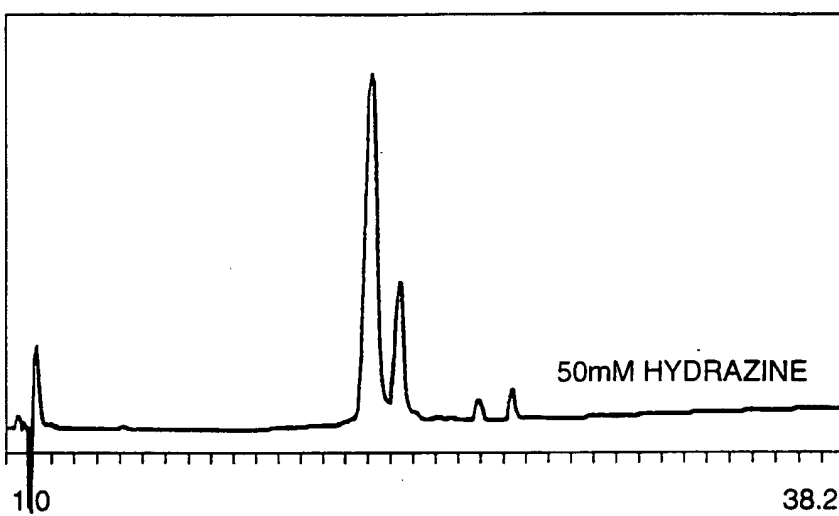


FIG._3B

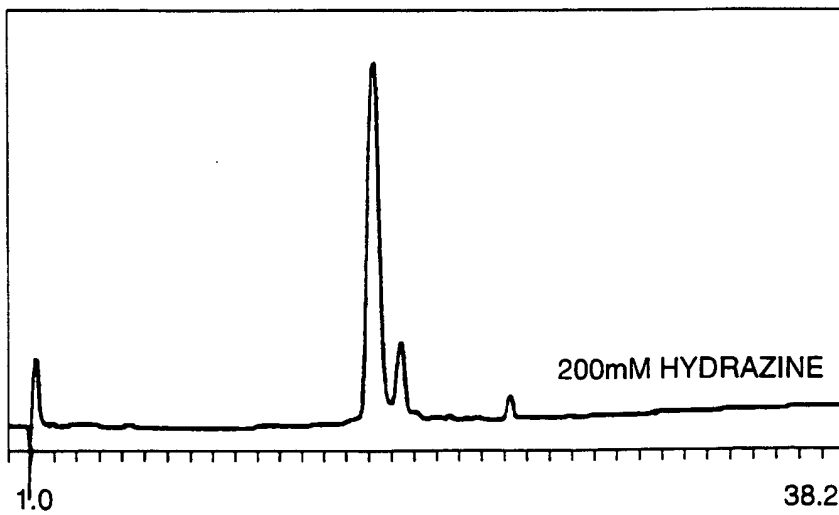


FIG._3C

4 / 31

Lev - MSC - LTEGLHGFHVHEFGDNTAGCTSAGPHFNPLSRKHG - COSH
MW=4022.4

*H - LTEGLHGTVHEFGDNTAGCTSAGPHFNPLSRKHG - COSH
MW=3745.1*

ALIQUOT OF PEPTIDE IN 6M Gu-HCl, 0.1M NaAc, pH 4.6 WAS DILUTED INTO 6M Gu-HCl, 0.1M NaAc, pH 14 FOR TWO MINUTES, QUENCHED WITH 6M Gu-HCl, 0.1M NaAc, pH 2.0

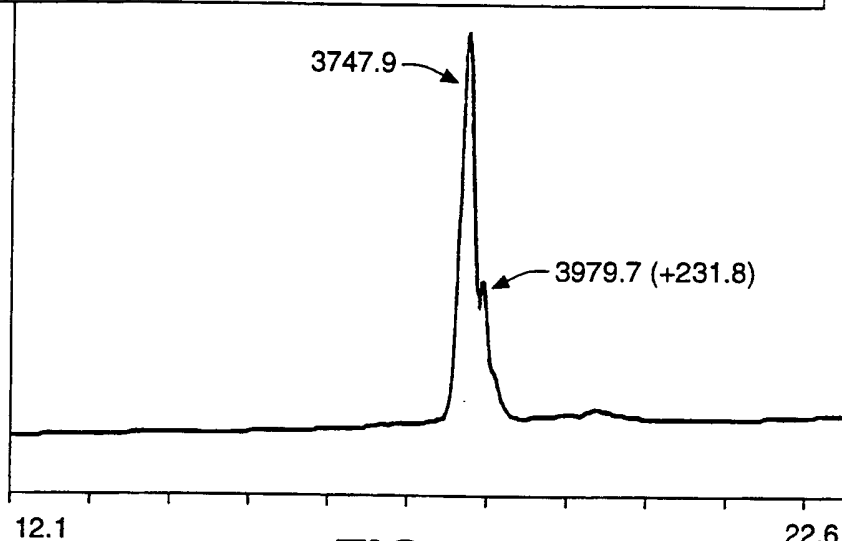


FIG.-4

1 21 47
TLQKKIEEIAAKYKHSVVKCCYDGACVNNDTCEQRAARISLGPKCIKAFTTECC
VVASQLRANISHKDMQLGR

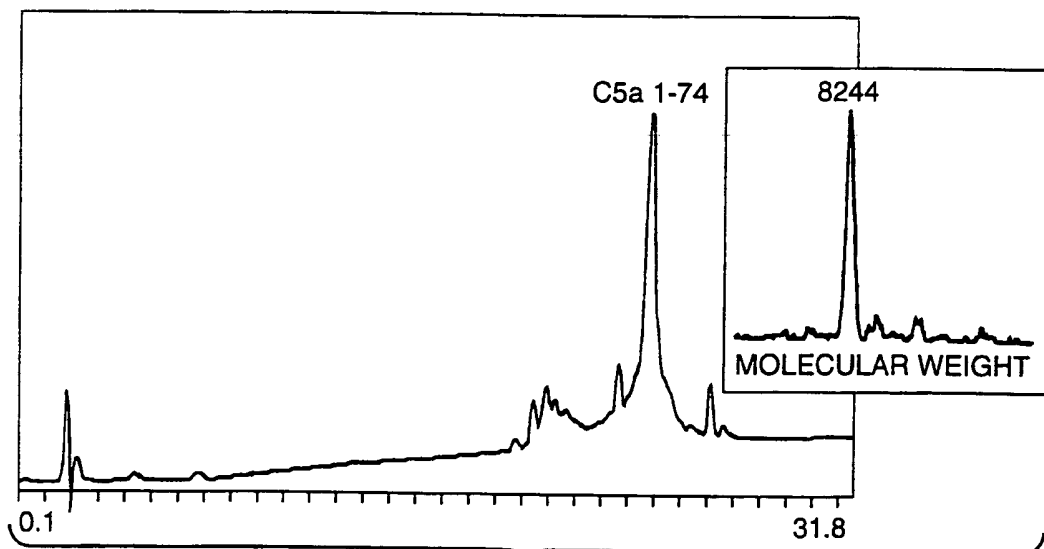
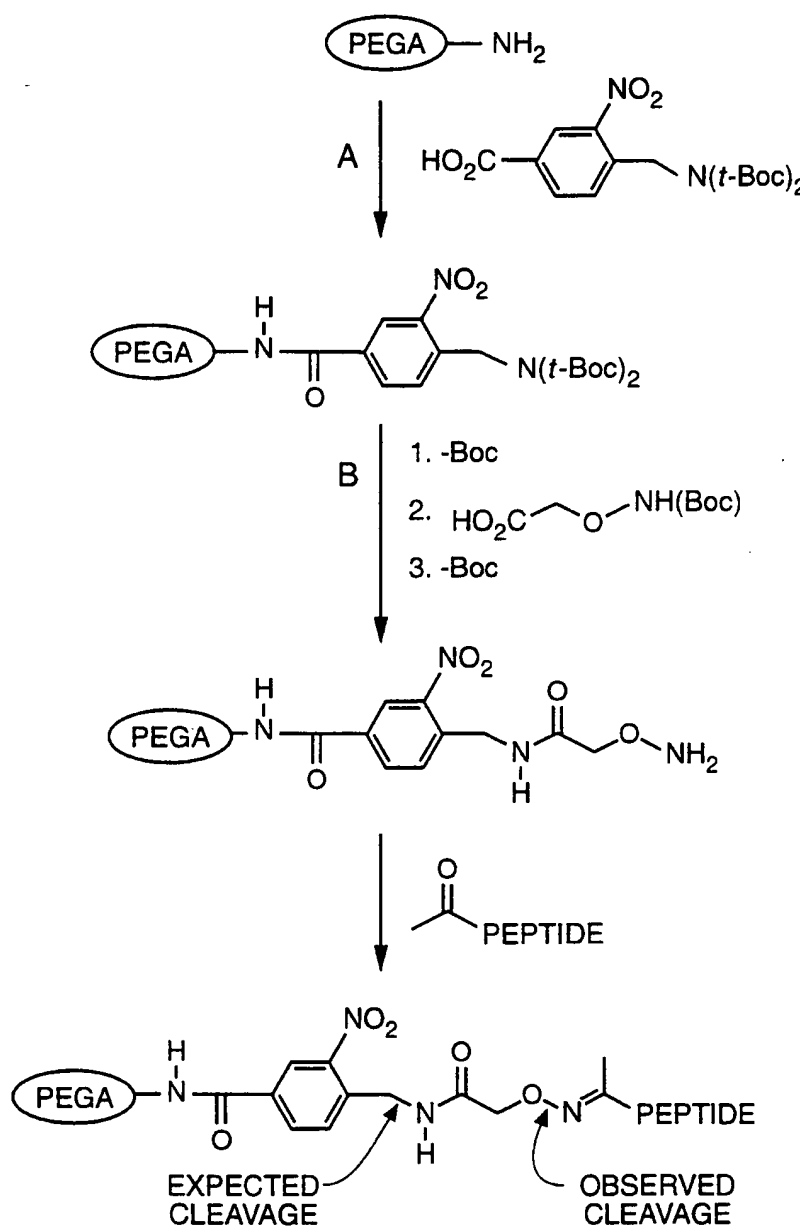
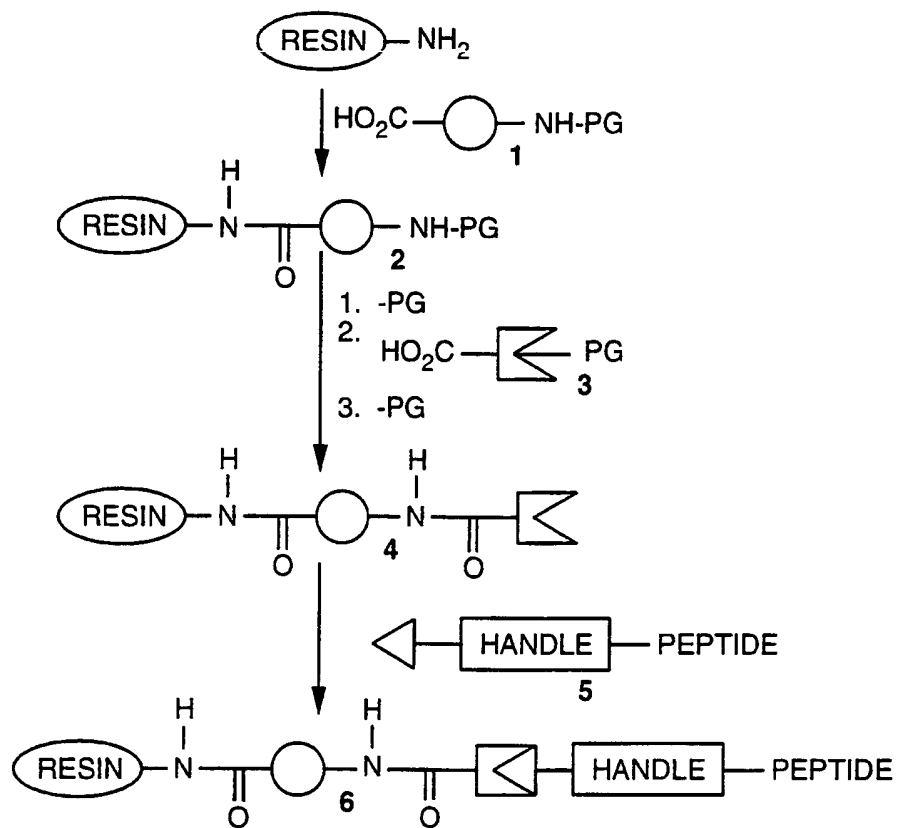


FIG.-26

5 / 31

**FIG.-5A**



$\text{HO}_2\text{C}-\text{O}-\text{NH}-$ = CLEAVABLE LINKER USED FOR MONITORING
WITH MALDI, ELECTROSPRAY MASS
SPECT, ETC...

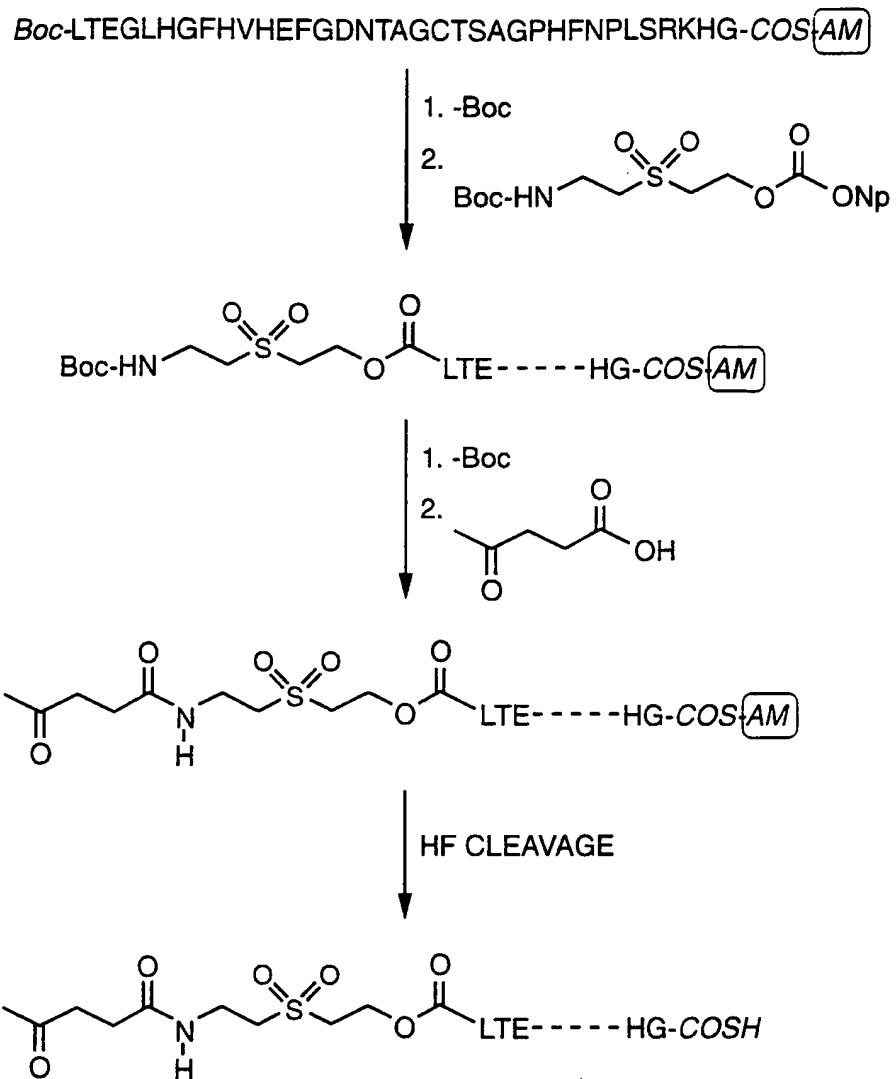
PG = PROTECTING GROUP

$\text{HO}_2\text{C}-\text{C}\backslash-$ = FUNCTIONAL GROUP ADDED TO RESIN TO
COUPLE WITH PEPTIDE

$\text{C}\backslash-\text{HANDLE}-\text{PEPTIDE}$ = PEPTIDE FUNCTIONALIZED WITH
1. CLEAVABLE HANDLE FOR RELEASE OF
PEPTIDE/PROTEIN FROM THE RESIN AT
COMPLETION OF SYNTHESIS AND
2. FUNCTIONAL GROUP TO COUPLE TO RESIN

FIG._5B

7 / 31

**FIG._6**

BEST AVAILABLE COPY

WO 98/56807

PCT/US98/12278

8 / 31

Lev - MSC - LTEGLHGFHVHEFGDNTAGCTSAGPHFNPLSRKHG - COSH (1)
+ Resin - PCL - ONH₂

↓ 1. pH 4.6, 6M Gu-HCl, 0.1 ACETATE

Resin - PCL - oxime - MSC - LTEGLHGFHVHEFGDNTAGCTSAGPHFNPLSRKHG - COSH (1)

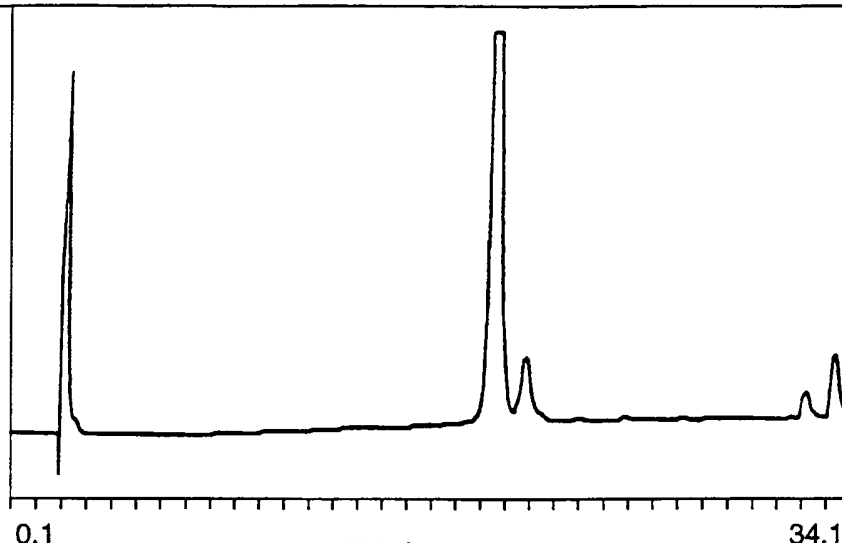


FIG._7A

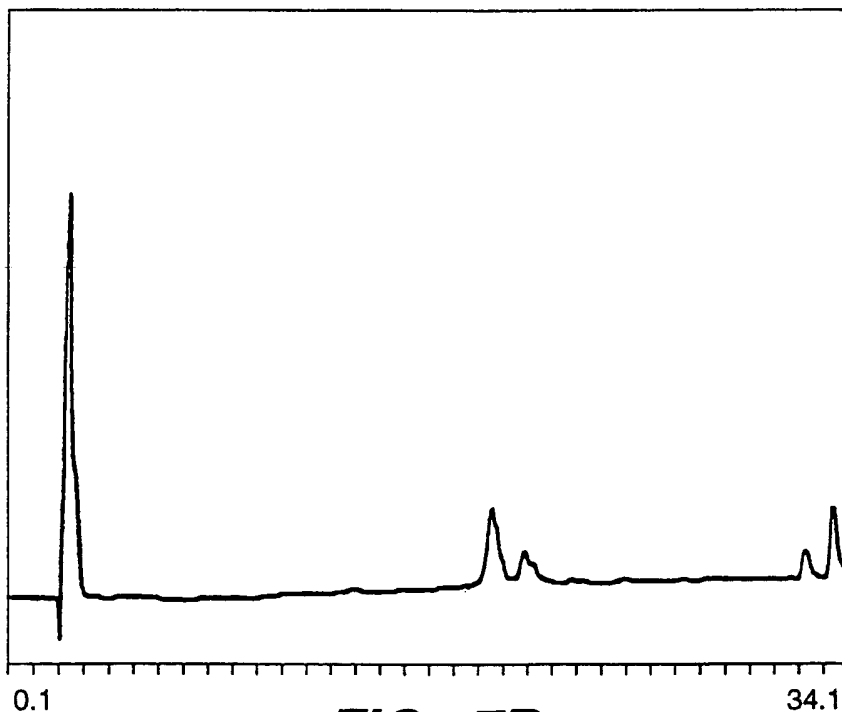


FIG._7B

BEST AVAILABLE COPY

WO 98/56807

PCT/US98/12278

9 / 31

Lev - MSC - LTEGLHGFHVHEFGDNTAGCTSAGPHFNPLSRKHG - COSH (1)
+ Resin - PCL - ONH₂

↓ 1. pH 4.6, 6M Gu•HCl, 0.1 ACETATE

Resin - PCL - oxime - MSC - LTEGLHGFHVHEFGDNTAGCTSAGPHFNPLSRKHG - COSH (1)

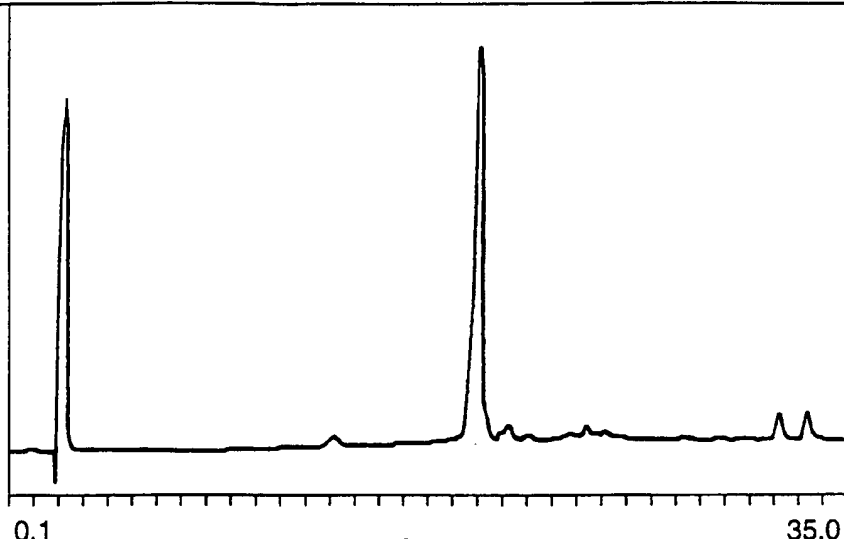


FIG._8A

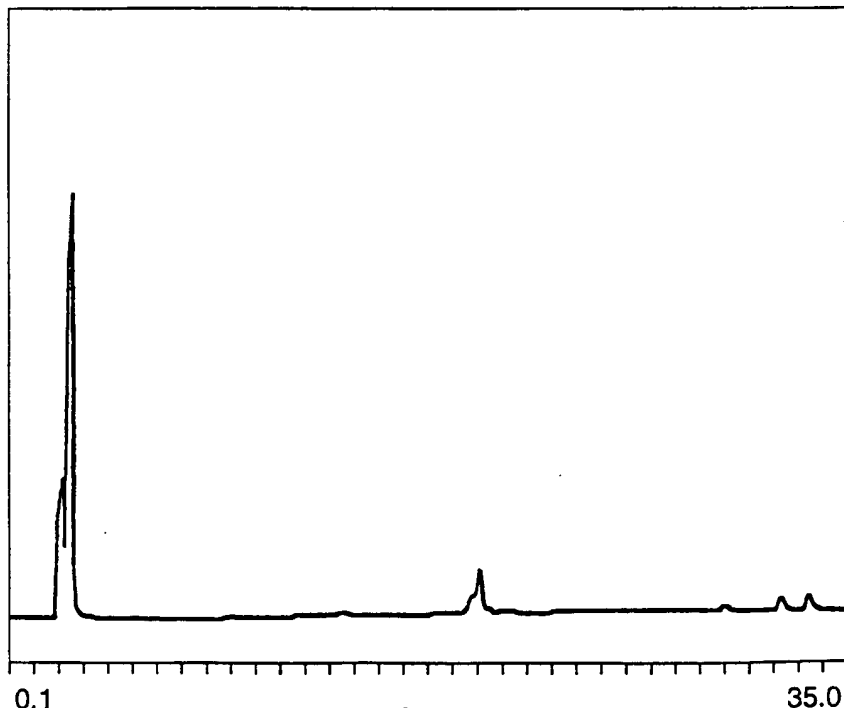


FIG._8B

10 / 31

Lev - MSC - LTEGLHGFHVHEFGDNTAGCTSAGPHFNPLSRKHG - COSH (1)
+ Resin - PCL - ONH2

↓ 1. pH 4.6, 6M Gu-HCl, 0.1 ACETATE

Resin - PCL - oxime - MSC - LTEGLHGFHVHEFGDNTAGCTSAGPHFNPLSRKHG - COSH (1)
MALDI MASS = 4022, BASE CLEAVAGE MASS = 3745

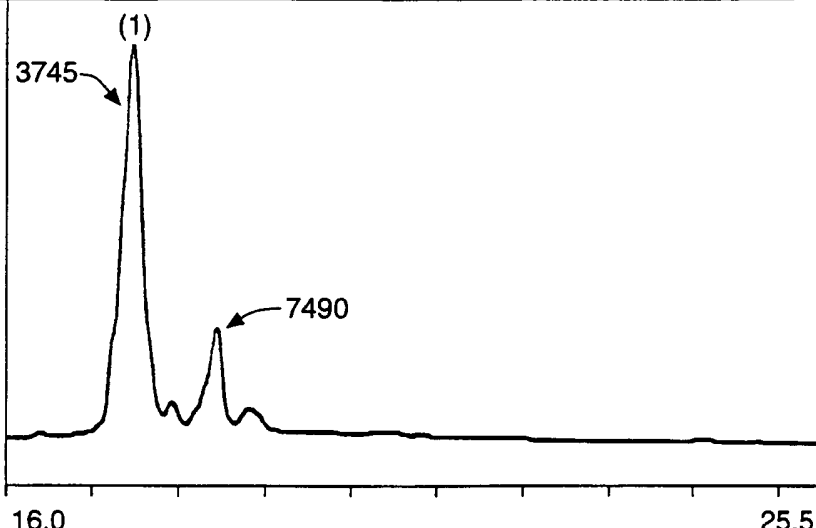


FIG._9A

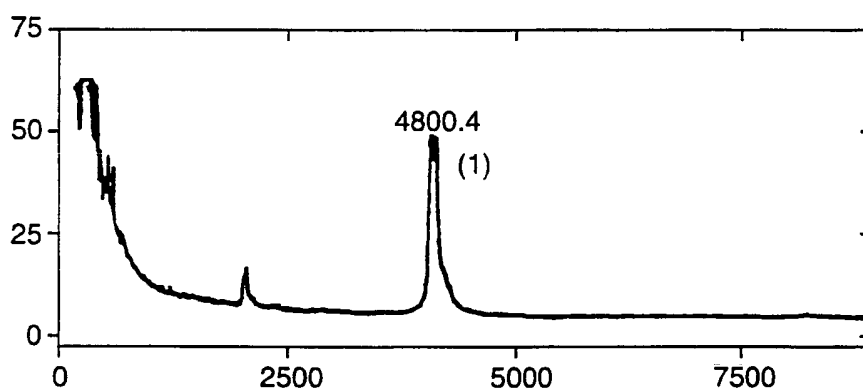


FIG._9B

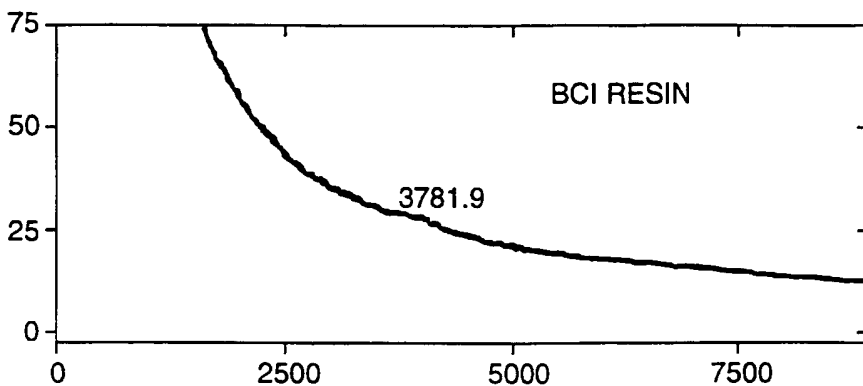


FIG._9C

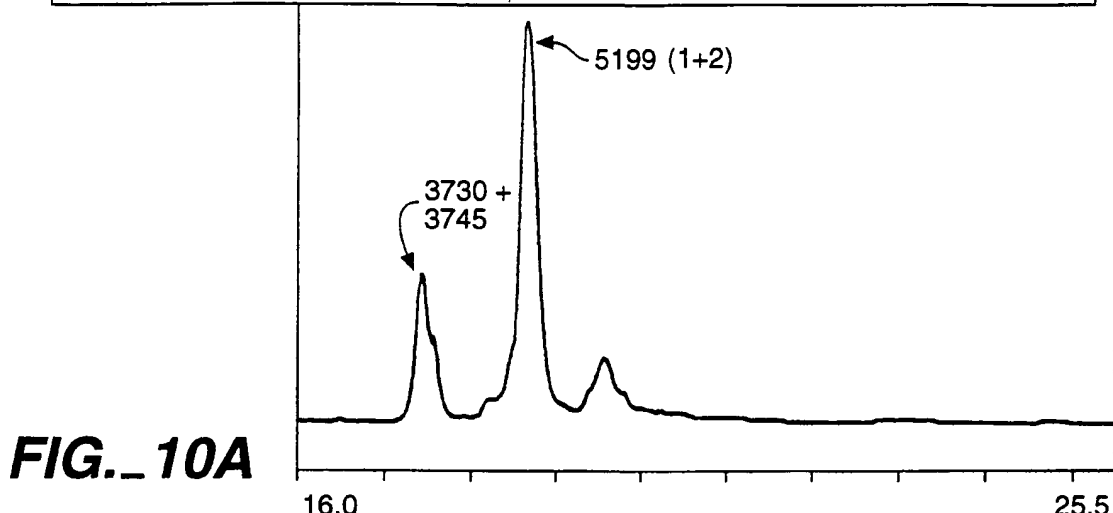
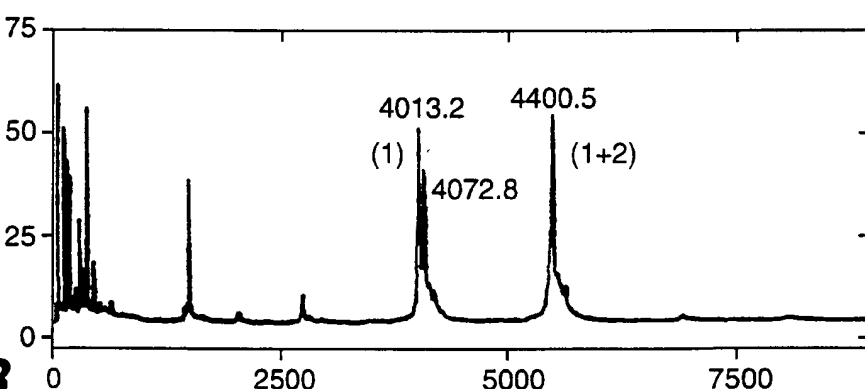
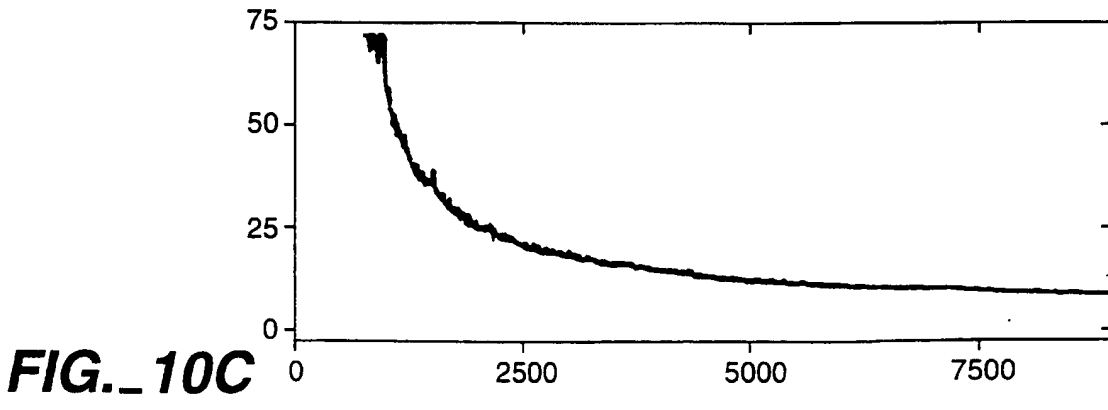
11 / 31

Resin - PCL - oxime - MSC - LTEGLHGFWHEFGDNTAGCTSAGPHFNPLSRKHG - COSAc (1)
MALDI MASS = 4080, BASE CLEAVAGE MASS = 3729
+ H - CGFRVREFGDNTA - COSH (2)

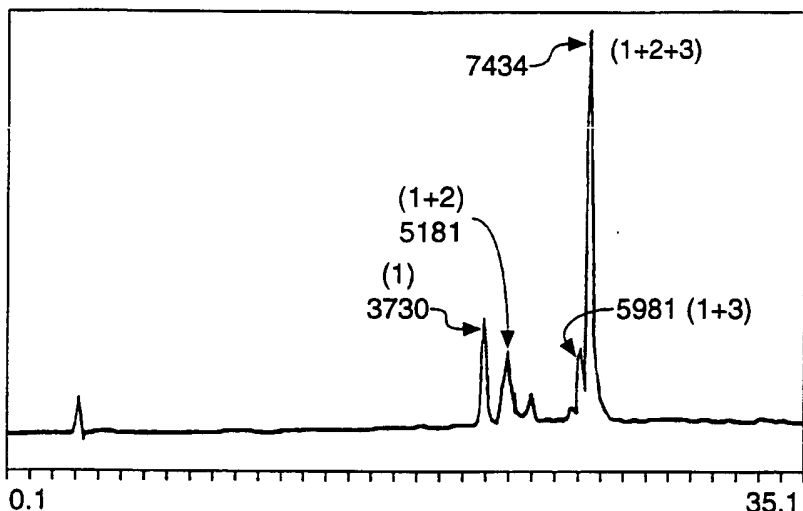
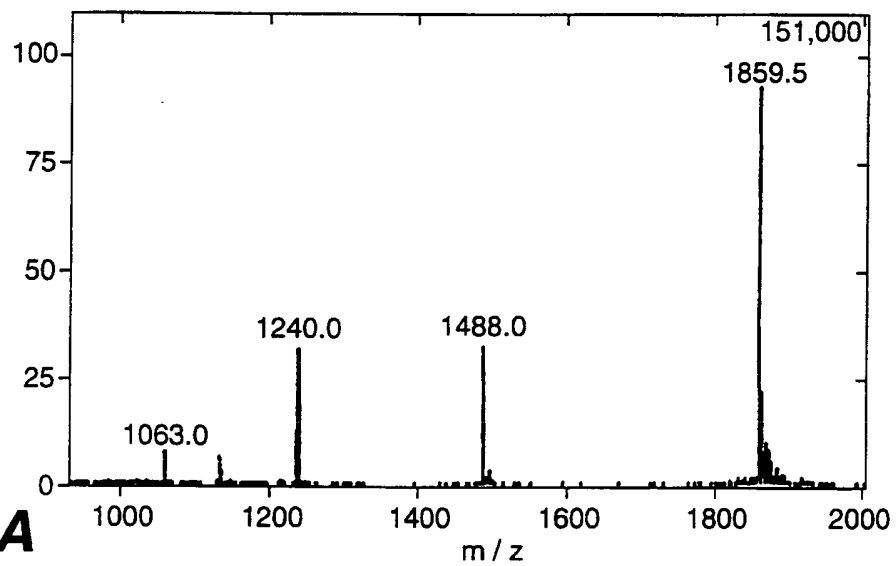
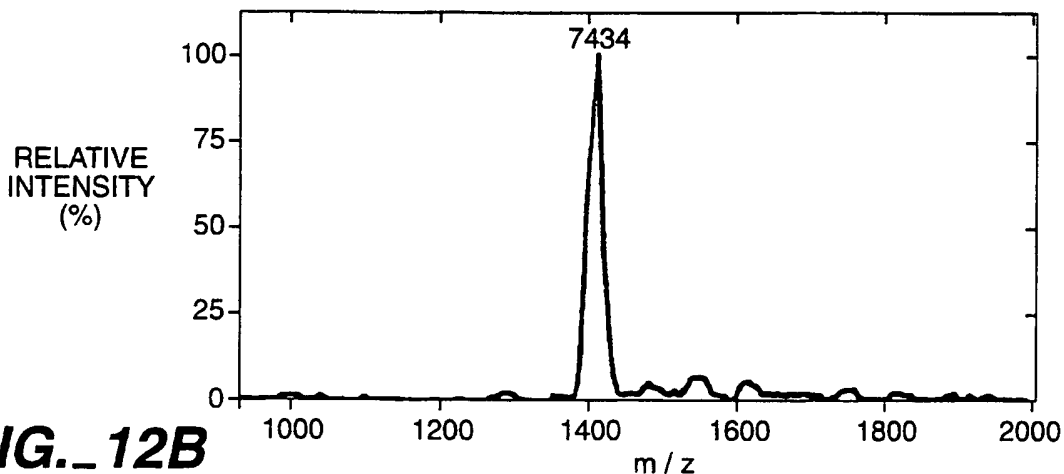
↓ 3. pH 7.5, 6M Gu•HCl, 0.1M PHOSPHATE, 0.5% THIOPHENOL

Resin - PCL - oxime - MSC - LTEGLHGFWHEFGDNTAGCTSAGPHFNPLSRKHGCGFRVREF -
GDNTA - COSH (1+2)

MALDI MASS = 5476, BASE CLEAVAGE MASS = 5199

**FIG._ 10A****FIG._ 10B****FIG._ 10C**

12 / 31

**FIG._ 11****FIG._ 12A****FIG._ 12B**

13 / 31

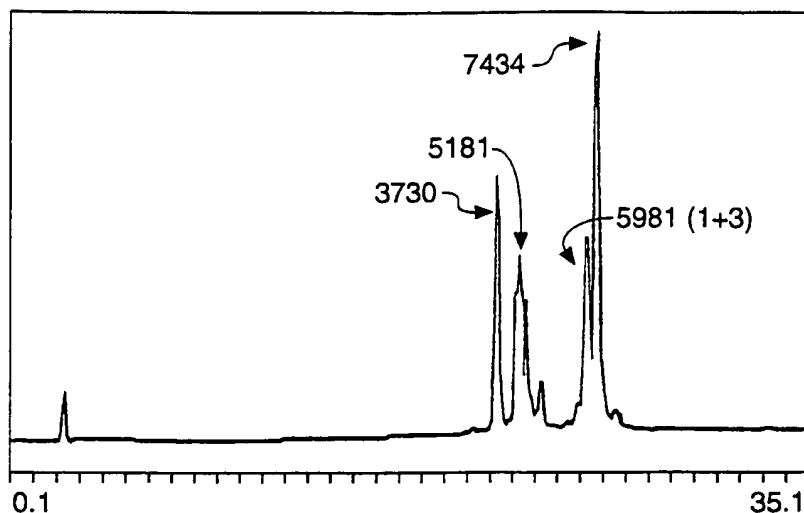


FIG._ 13

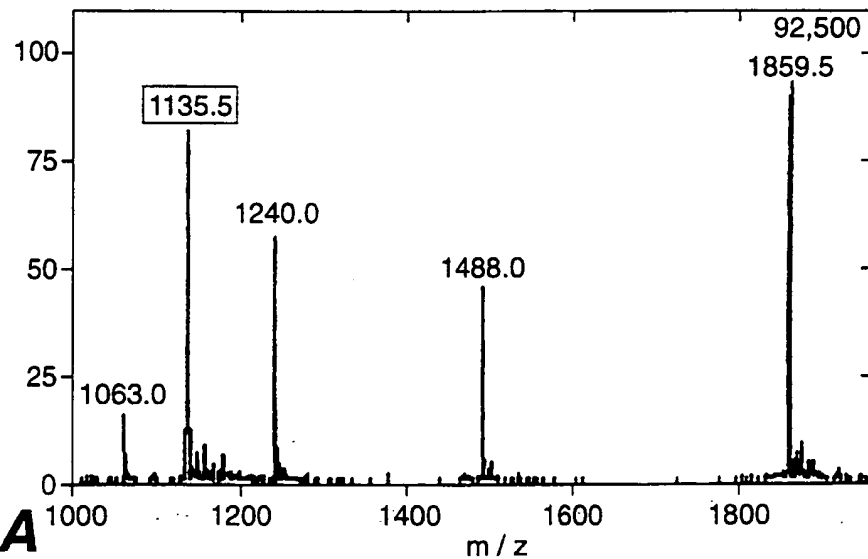


FIG._ 14A

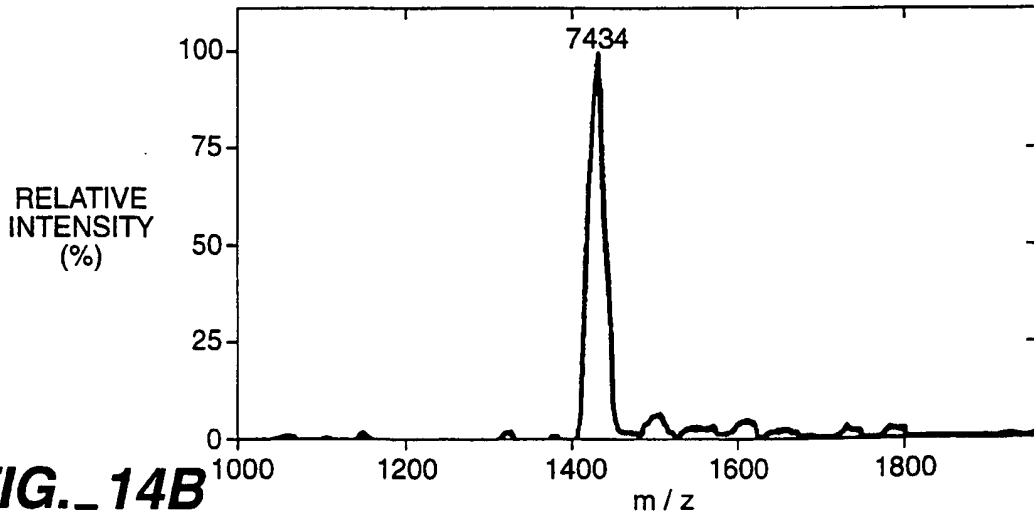


FIG._ 14B

BEST AVAILABLE COPY

WO 98/56807

PCT/US98/12278

14 / 31

FIG._ 15A

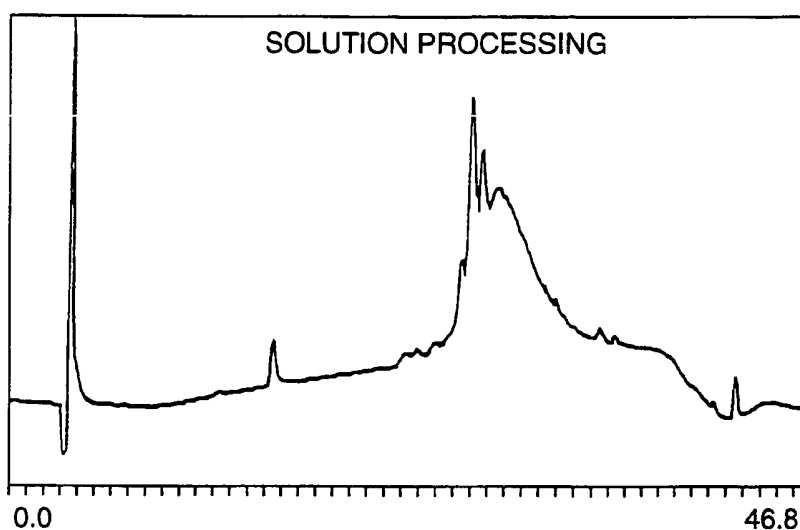


FIG._ 15B

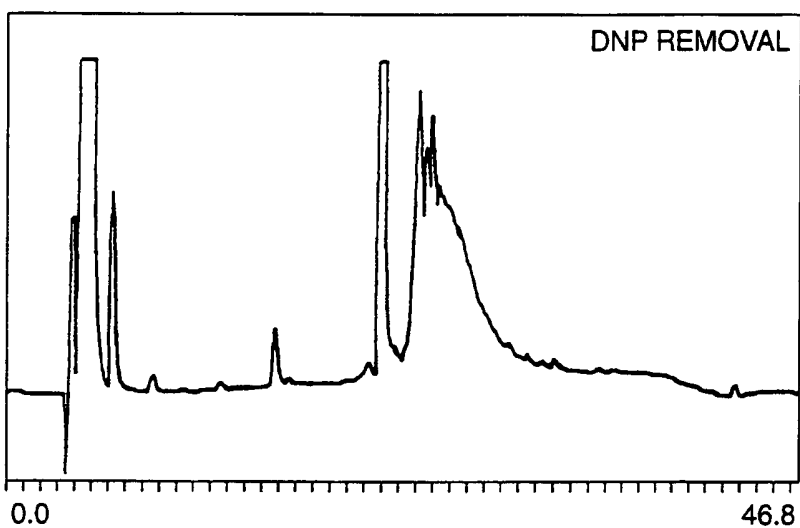
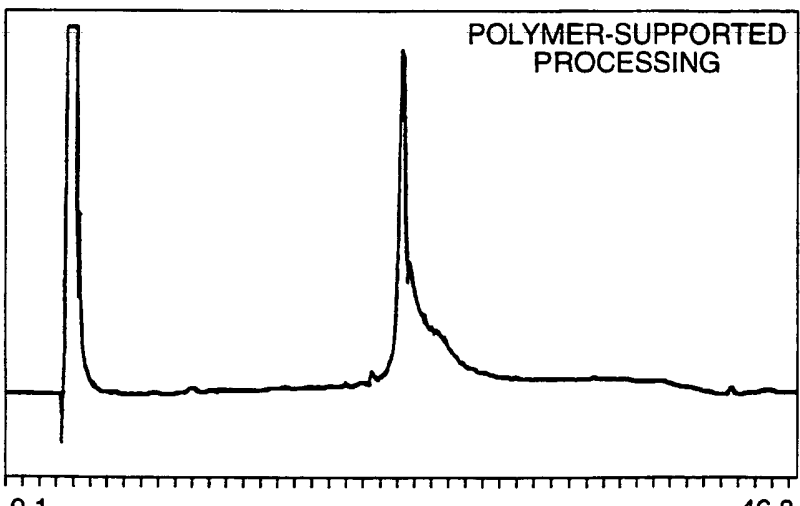


FIG._ 15C



SUBSTITUTE SHEET (RULE 26)

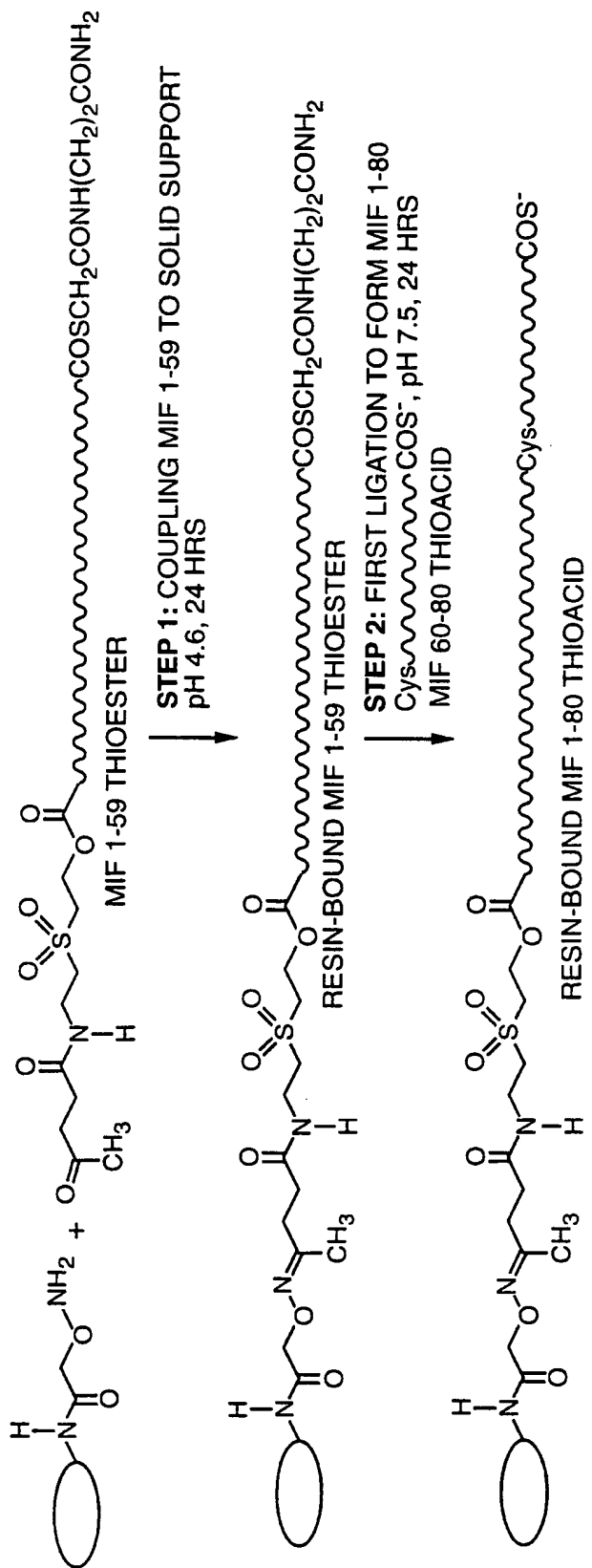
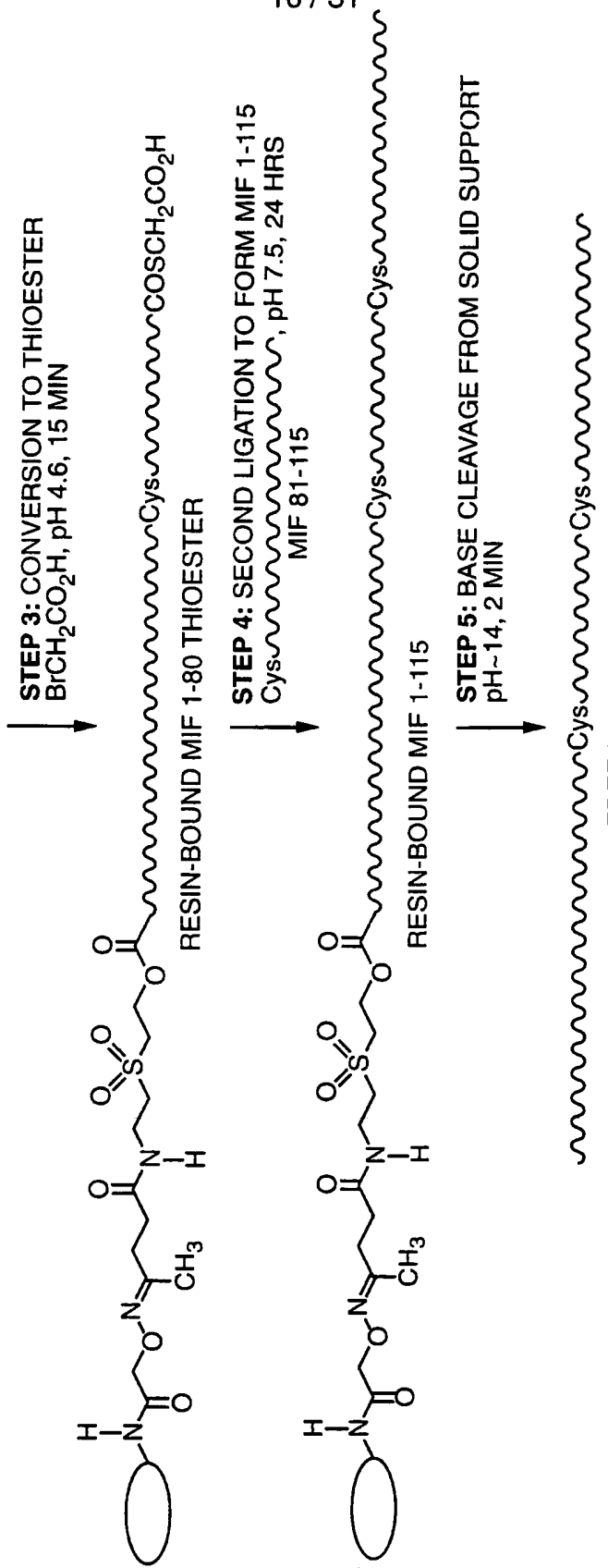
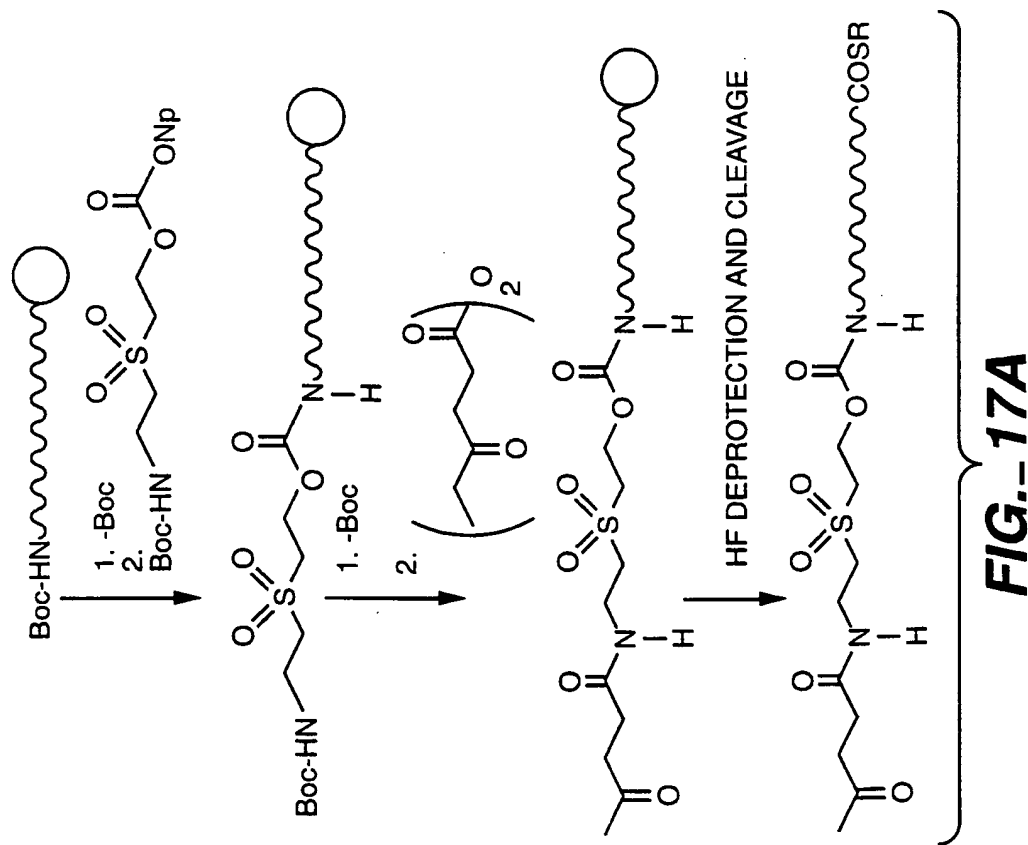
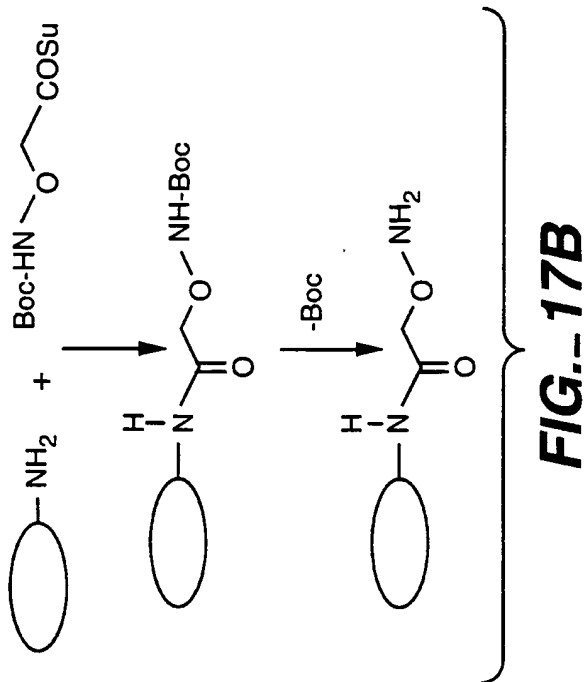


FIG. 16A

16 / 31

**FIG._ 16B**

17 / 31

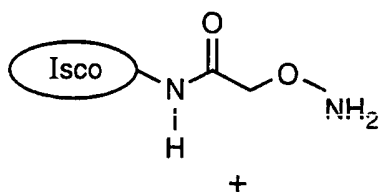


BEST AVAILABLE COPY

WO 98/56807

PCT/US98/12278

18 / 31



↓

Isco — OXIME - MSC HANDLE - MET¹ - MIF 2 - 58 - Leu⁵⁹ - SAc - βAla - CO₂H
EXPECTED BASE CLEAVAGE MASS = 6271

FIG._ 18A

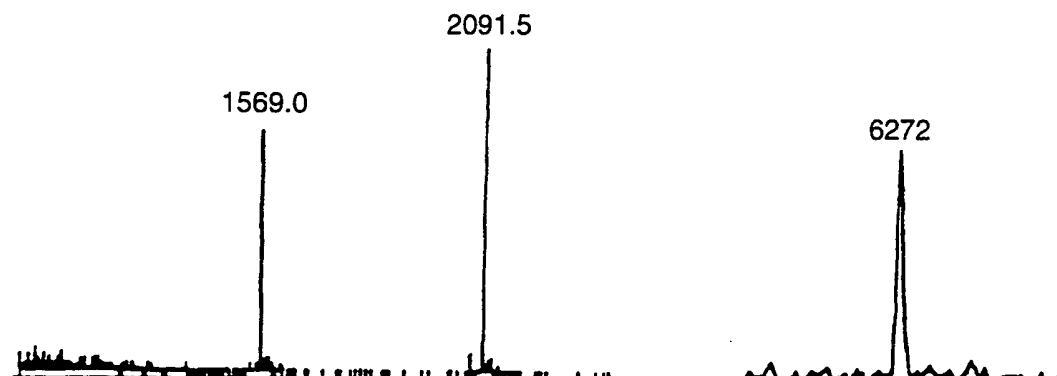


FIG._ 18C

FIG._ 18D

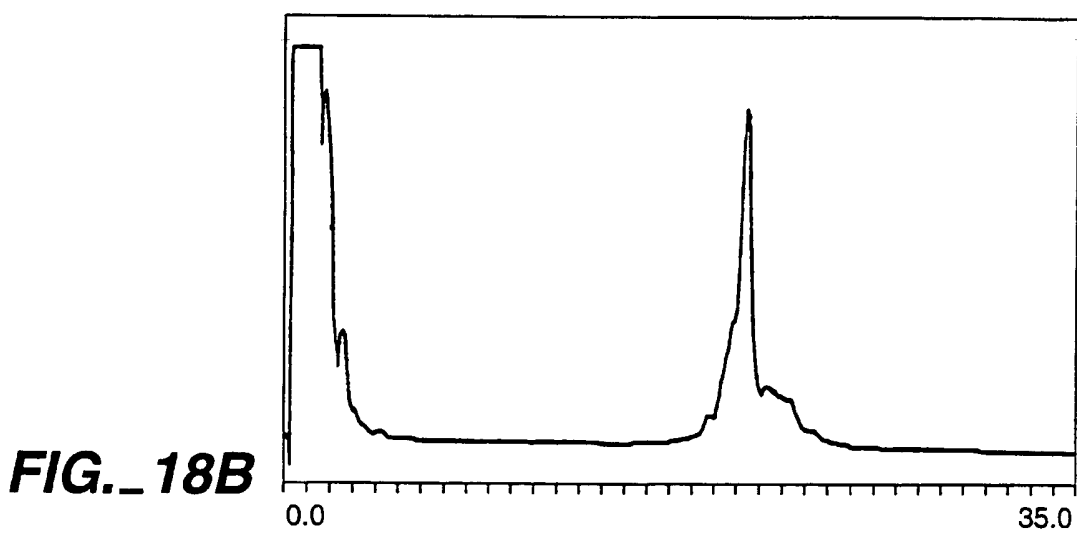


FIG._ 18B

BEST AVAILABLE COPY

WO 98/56807

PCT/US98/12278

19 / 31

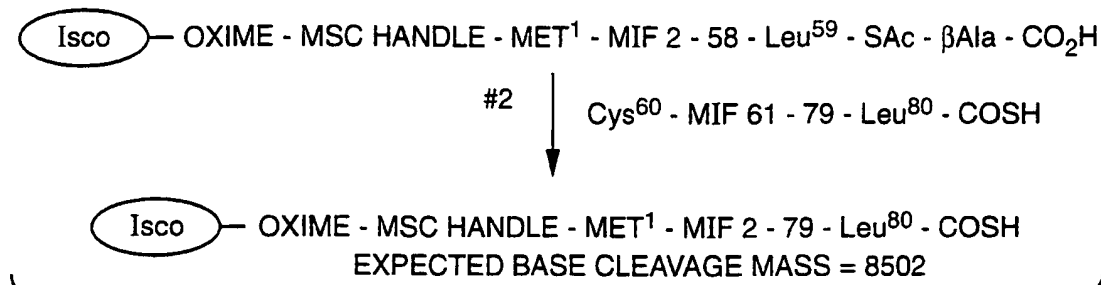


FIG. 19A

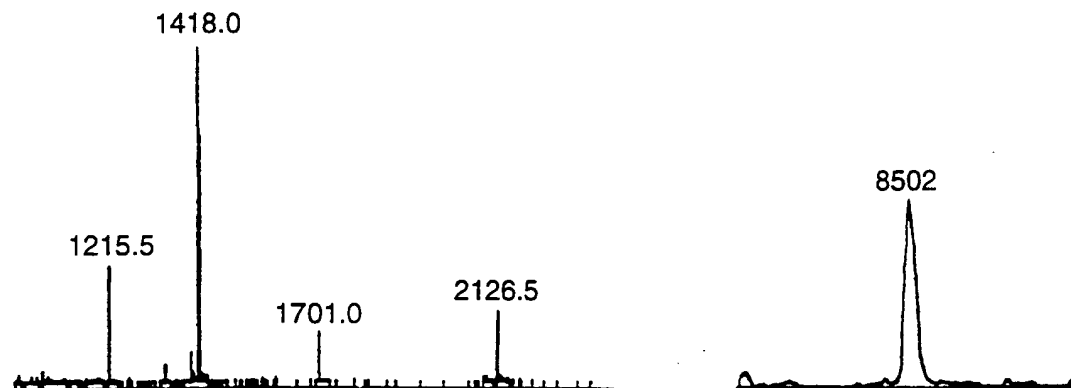


FIG. - 19C

FIG.-19D

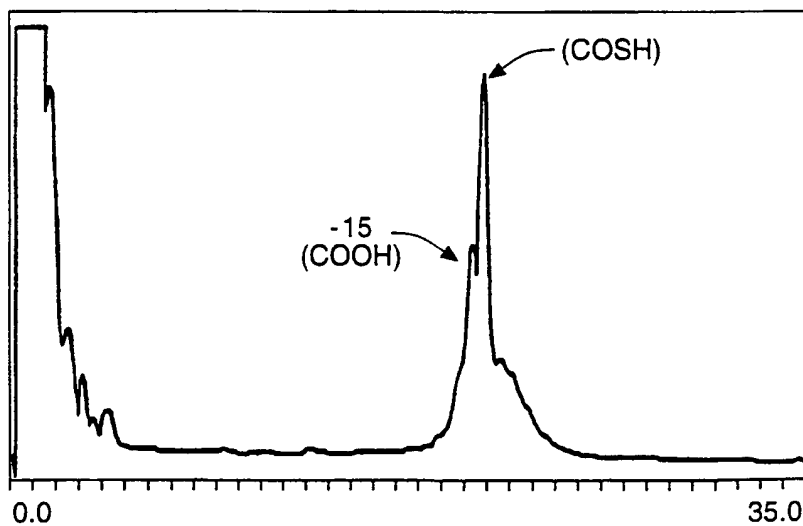
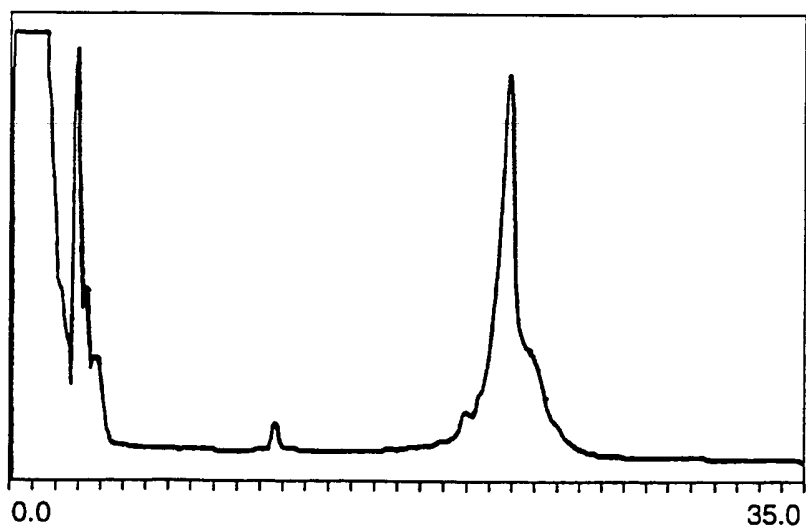
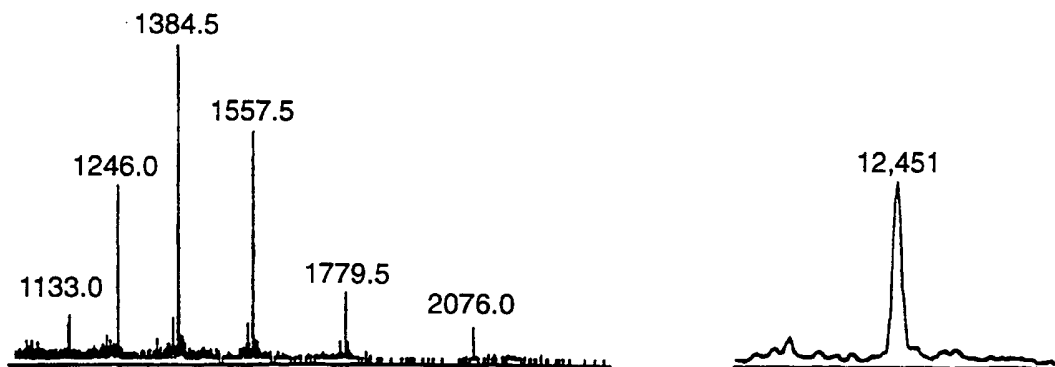
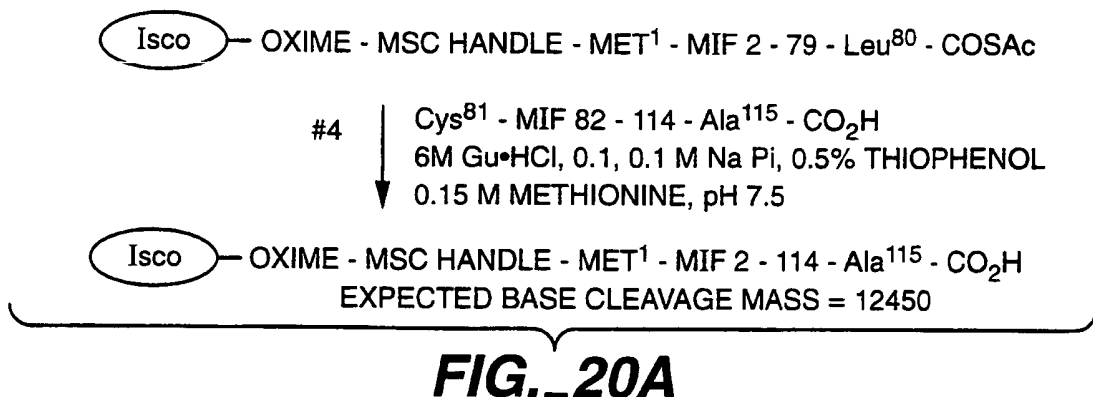
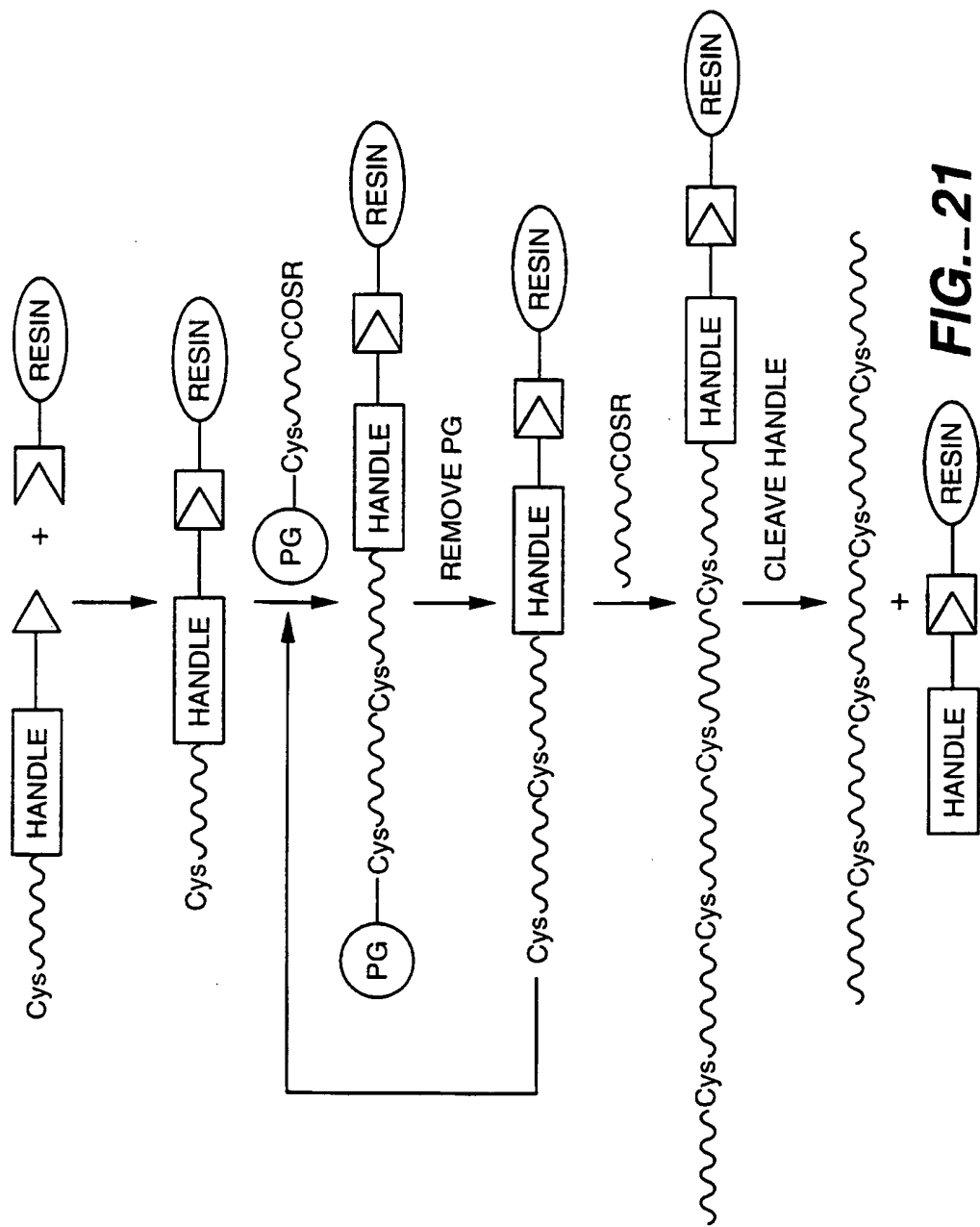
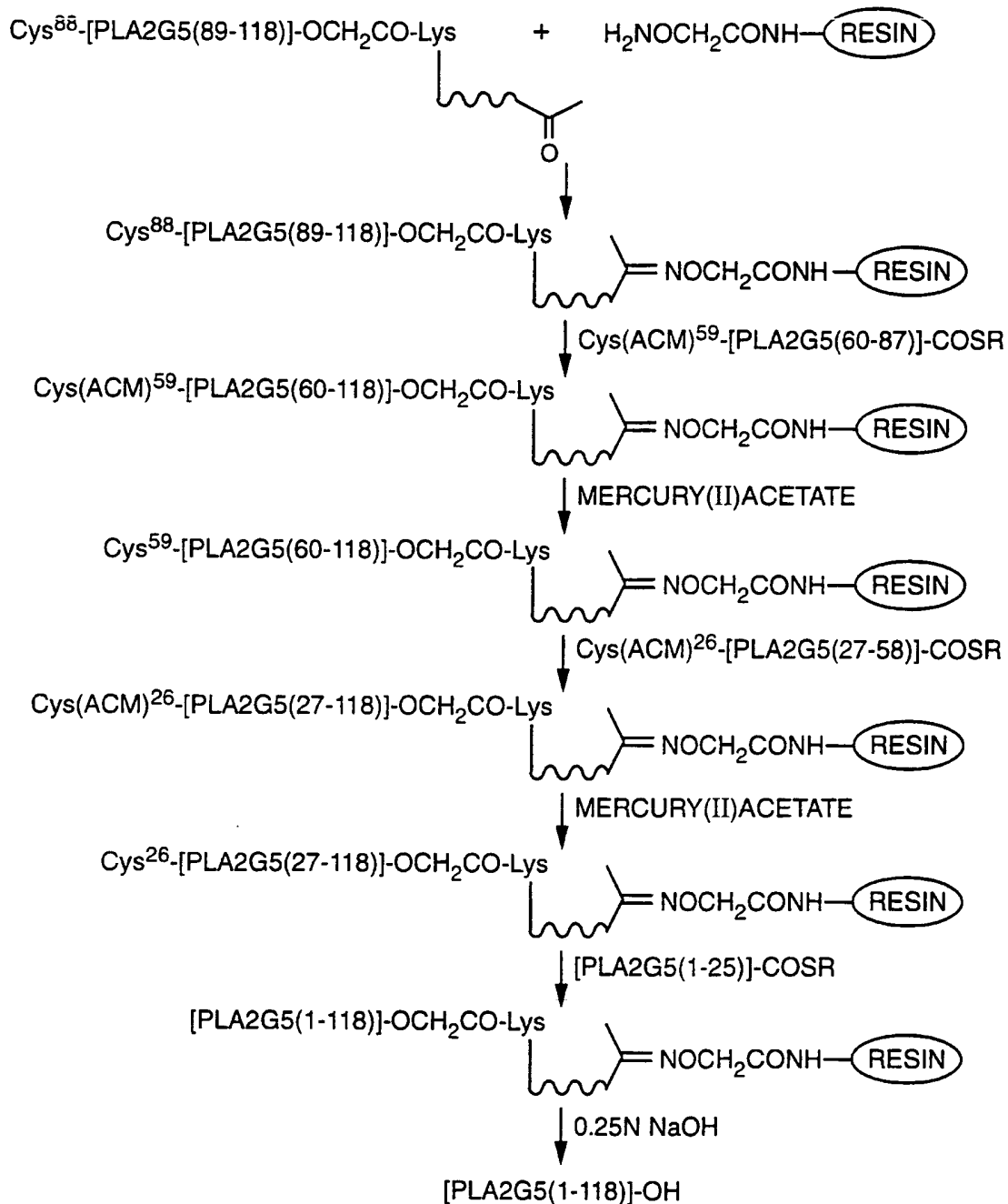


FIG._19B

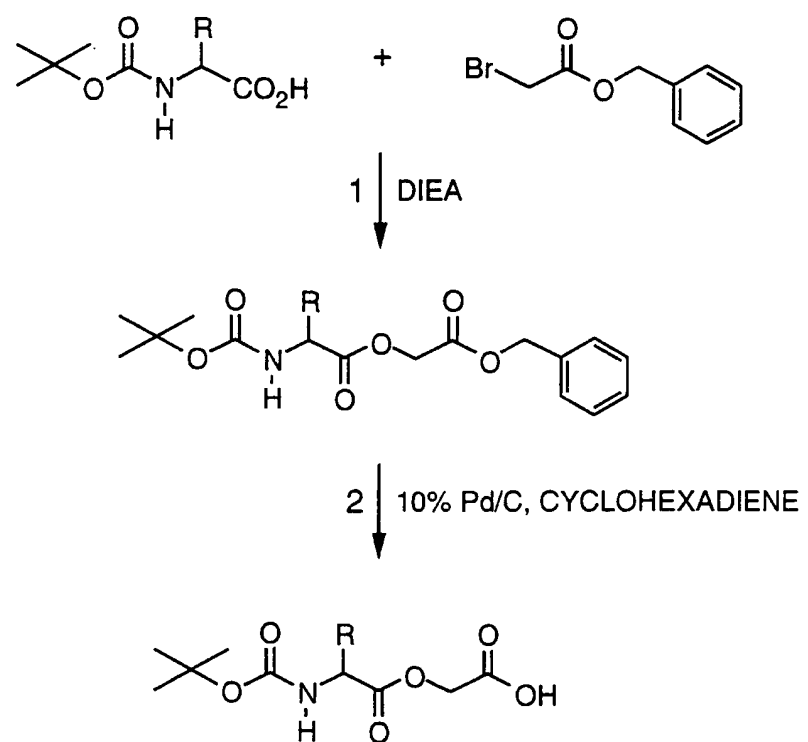
20 / 31

**FIG._20B**

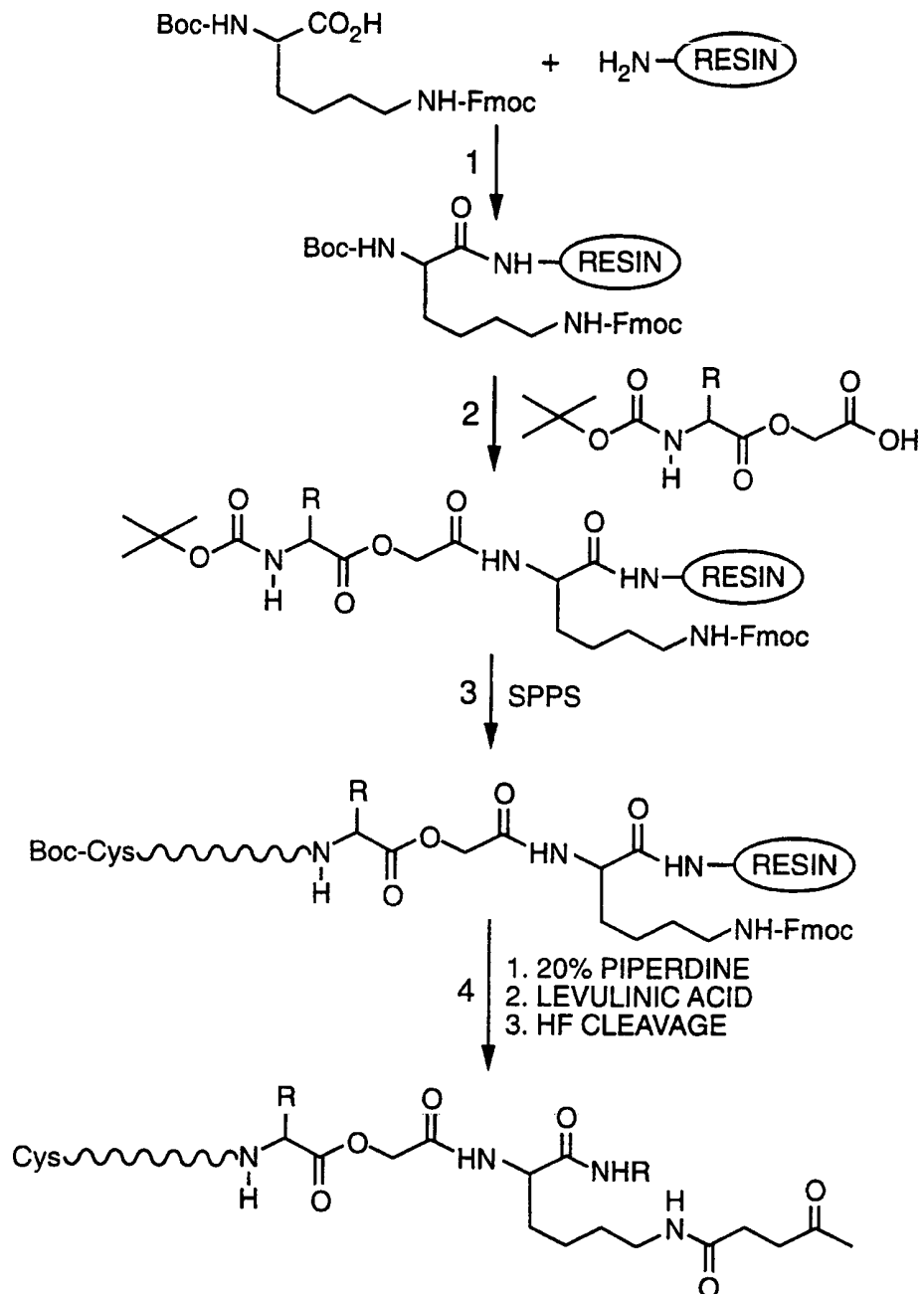


**FIG._22**

23 / 31

**FIG.-23**

24 / 31

***FIG._24***

25 / 31

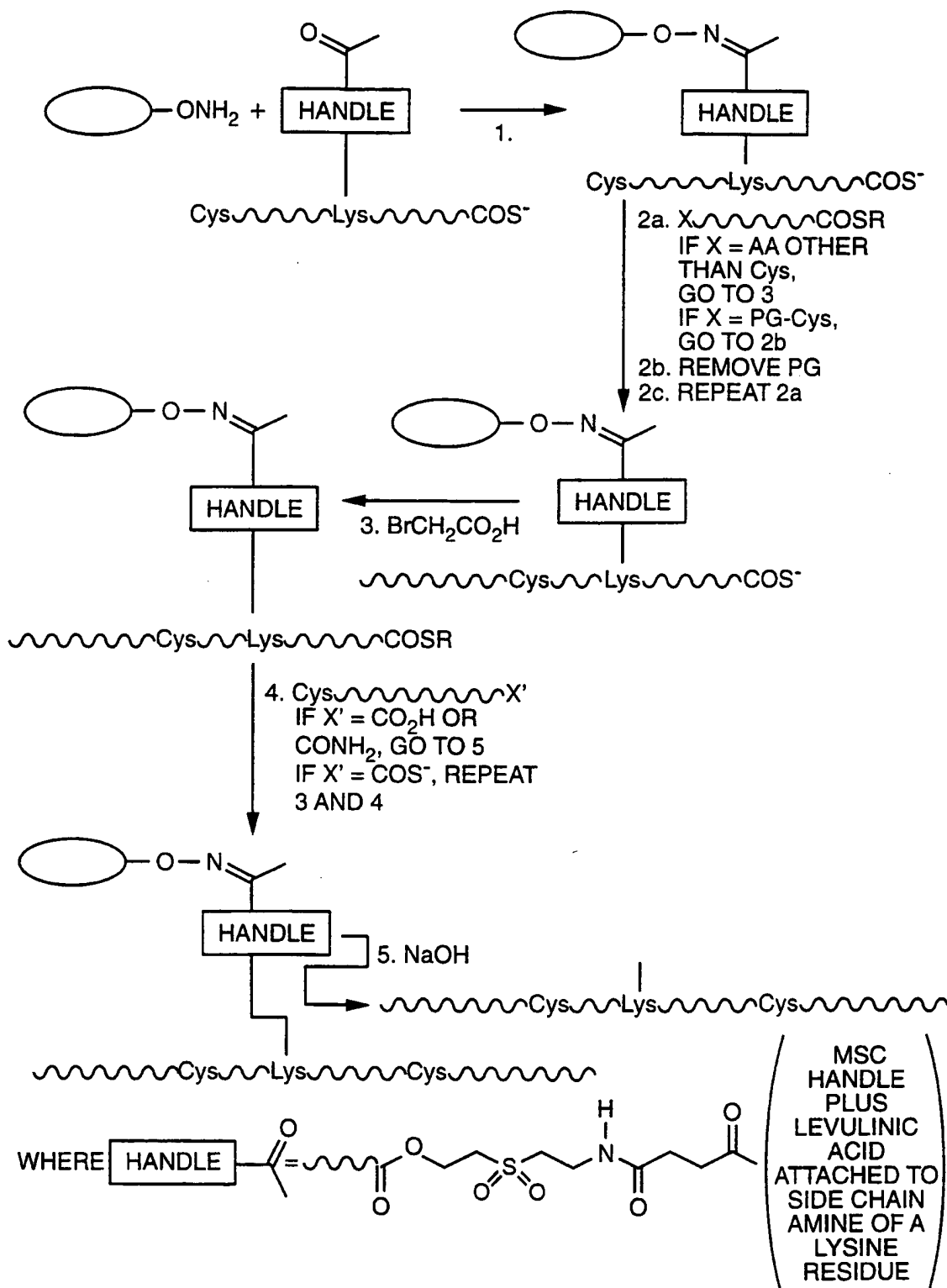
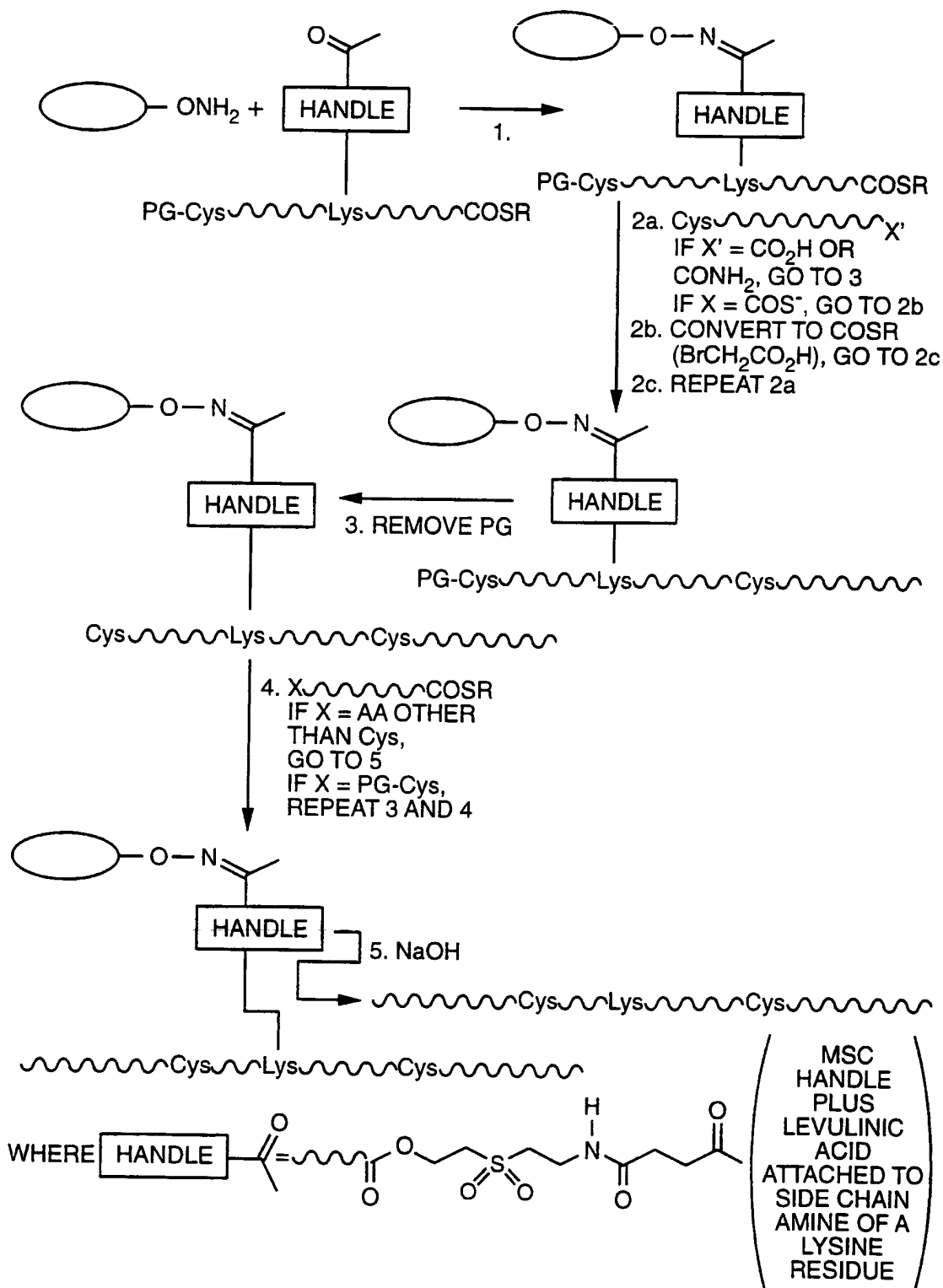


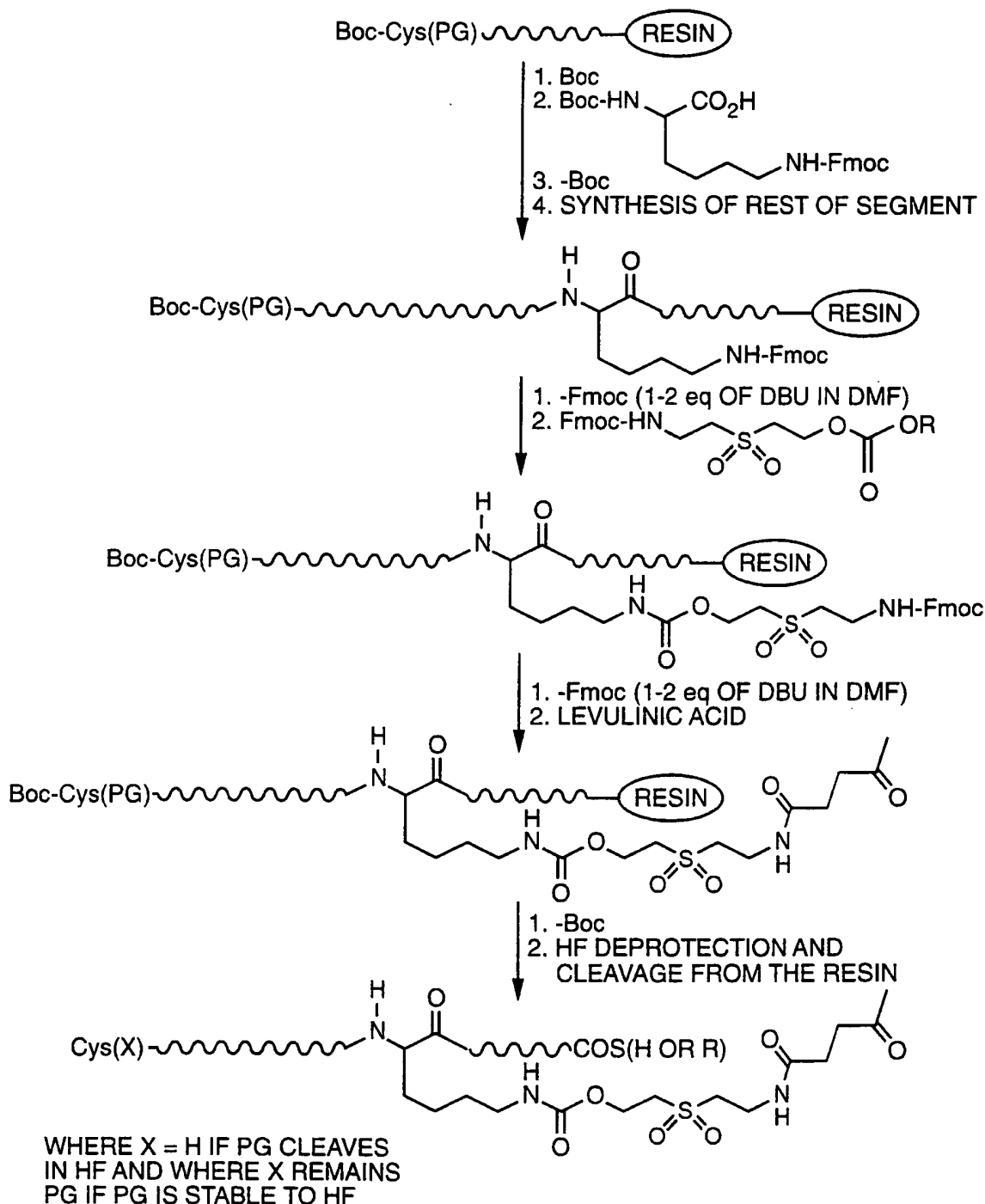
FIG.-25A

SUBSTITUTE SHEET (RULE 26)

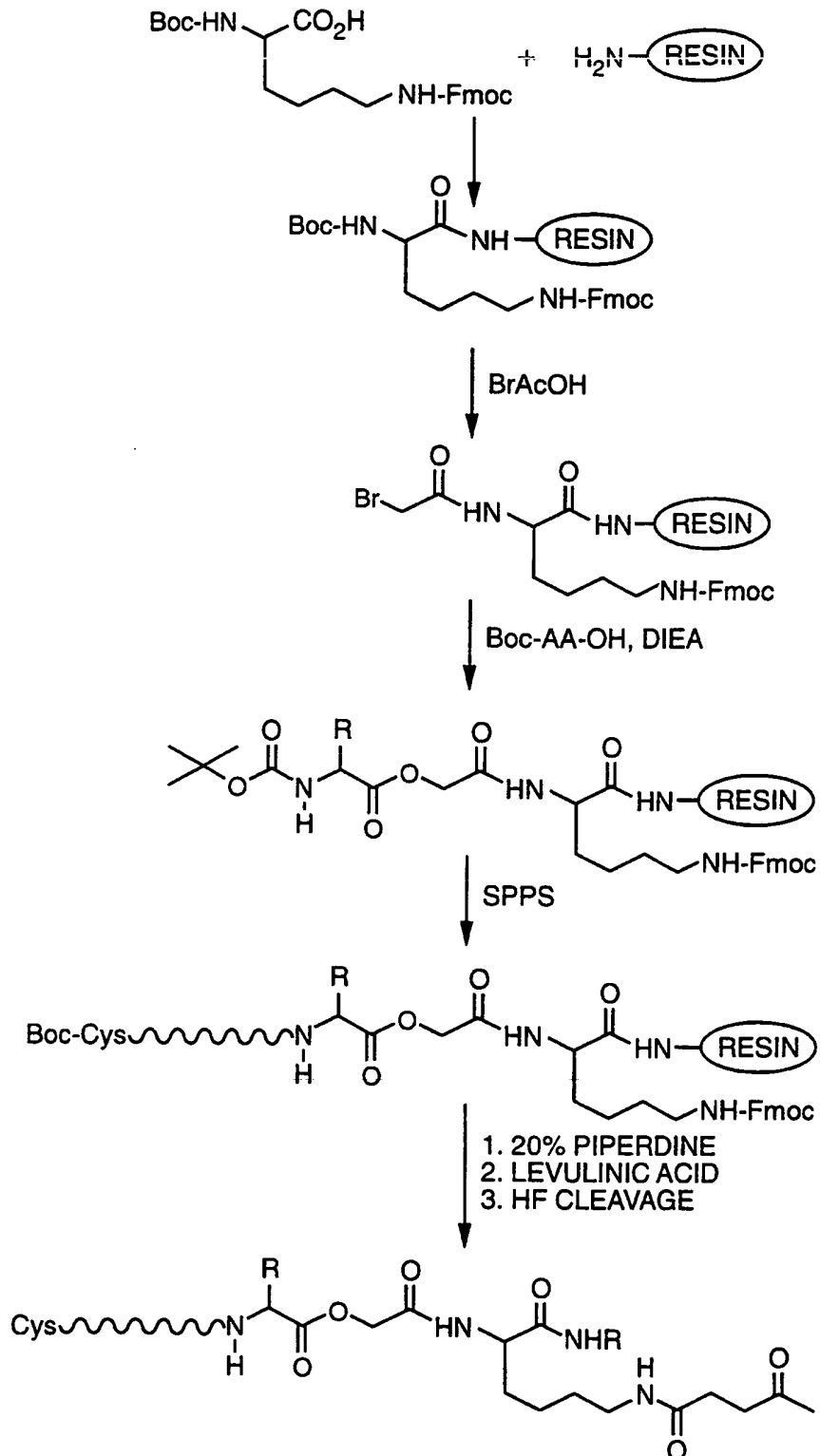
**FIG.-25B**

SUBSTITUTE SHEET (RULE 26)

27 / 31

***FIG._25C***

28 / 31

**FIG._27**

SUBSTITUTE SHEET (RULE 26)

BEST AVAILABLE COPY

WO 98/56807

PCT/US98/12278

29 / 31

ALTKYGFYGCYGRLEEKGCADRKNILA
1 10 19 27

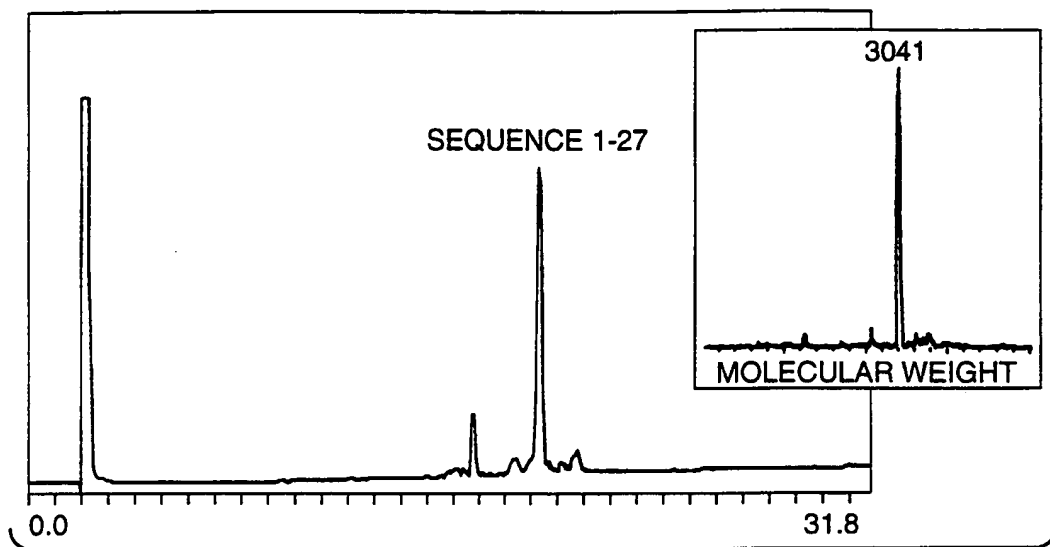


FIG._28

ALTKYGFYGCYGRLEEKGCADRKNILA
1 10 19 27

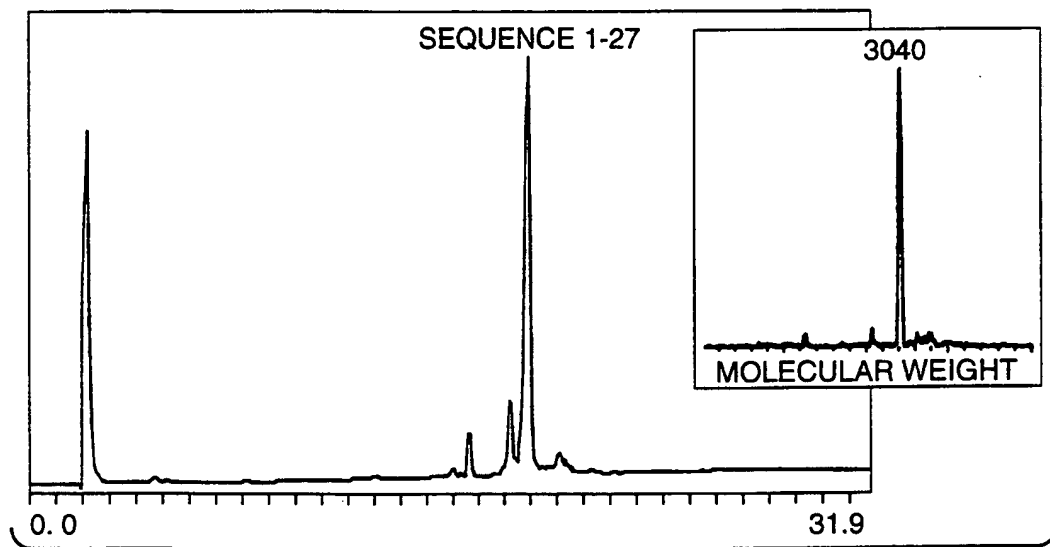
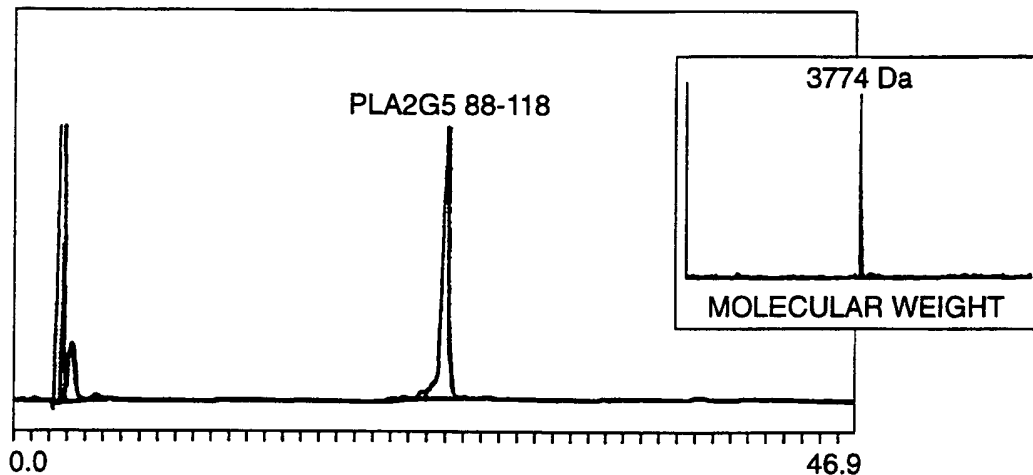


FIG._29

30 / 31

1	26	59
GLLDLKSMIEKVTKNALTNYGFYGCYCGWGGRGTPKDGTDWCCWAHDHCYGRLEEKGC NIRTQSYKYRFAWGVTCEPGPFCHVNLCACDRKLVYCLKRNLRSYNPQYQYFPNILCS		
88		118



1 26 59
GLLDLKSMIEKVTKNALTNYGFYGCYCGWGGRGTPKDGTDWCCWAHDHCYGRLEEKGC
NIRTQSYKYRFAWGVTCEPGPFCHVNLACACDRKLVYCLKRNLRSPNPQYQYFPNILCS
88 118

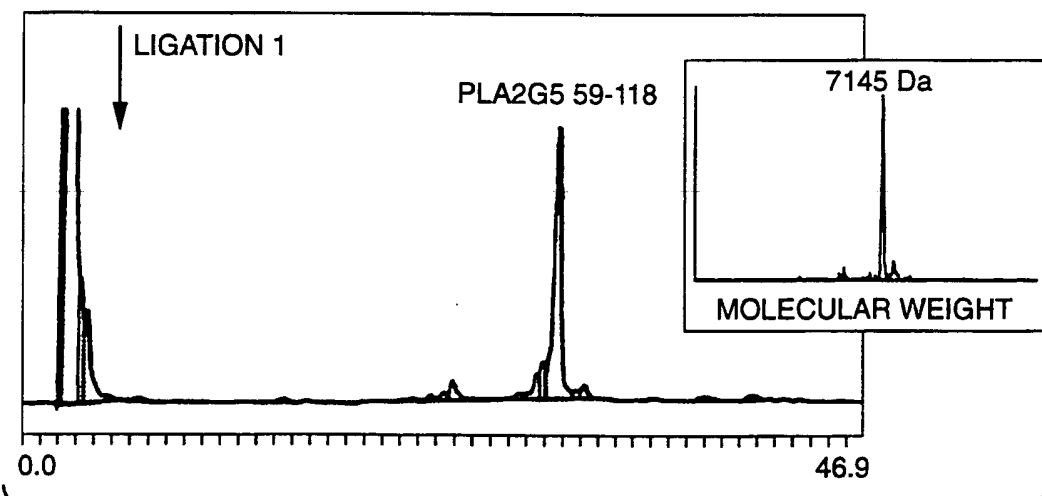


FIG.-30A

31 / 31

Mass spectrum showing the molecular weight distribution of PLA2G5 26-118. The main plot displays peaks across a range of molecular weights, with a prominent peak at 10876 Da. An inset provides a detailed view of the region around this peak. The x-axis is labeled "MOLECULAR WEIGHT" and includes numerical markers at 0.0 and 46.9. A vertical arrow labeled "LIGATION 2" points to the first two peaks on the left side of the main plot.

1 26 59

GLLDLKSMIEKVGTGNALTNYGFYGCYCGWGGRGTPKDGTDWCCWAHDHCYGRLEEKGC
NIRTSQSYKYRFAWGVVTCEPGPFCHVNLCACDRKLVYCLKRNLRSYNPQYQYFPNILCS

88 118

LIGATION 3

PLA2G5 1-118

13593 Da

MOLECULAR WEIGHT

0.0 46.9

FIG.-30B

FIG.-30A

FIG._30

INTERNATIONAL SEARCH REPORT

International Application No
PCT/US 98/12278

A. CLASSIFICATION OF SUBJECT MATTER
IPC 6 C07K1/02 C07K1/04 C07K1/08

According to International Patent Classification(IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)
IPC 6 C07K

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	WO 96 34878 A (SCRIPPS) 7 November 1996 cited in the application see the whole document ---	1-28
A	H HOJO & S AIMOTO: "Polypeptide synthesis using the S-alkyl thioester of a partially protected peptide segment. Synthesis of DNA-binding domain of c-myb protein (142-193)-NH2." BULLETIN OF THE CHEMICAL SOCIETY OF JAPAN., vol. 64, no. 1, January 1991, pages 111-117, XP002080326 TOKYO JP see the whole document ---	1-28 -/-

Further documents are listed in the continuation of box C.

Patent family members are listed in annex.

* Special categories of cited documents :

- "A" document defining the general state of the art which is not considered to be of particular relevance
- "E" earlier document but published on or after the international filing date
- "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
- "O" document referring to an oral disclosure, use, exhibition or other means
- "P" document published prior to the international filing date but later than the priority date claimed

- "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
- "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
- "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.
- "&" document member of the same patent family

Date of the actual completion of the international search

12 October 1998

Date of mailing of the International search report

27/10/1998

Name and mailing address of the ISA
European Patent Office, P.B. 5818 Patentlaan 2
NL - 2280 HV Rijswijk
Tel. (+31-70) 340-2040, Tx. 31 651 epo nl
Fax: (+31-70) 340-3016

Authorized officer

Masturzo, P

INTERNATIONAL SEARCH REPORT

International Application No
PCT/US 98/12278

C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT		
Category	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	H HOJO ET AL.: "Development of a linker with an enhanced stability for the preparation of peptide thioesters and its application to the synthesis of a stable-isotope-labelled HU-type DNA-binding protein." BULLETIN OF THE CHEMICAL SOCIETY OF JAPAN., vol. 66, no. 9, September 1993, pages 2700-2706, XP002080327 TOKYO JP see the whole document ---	1-28
A	CHEMICAL ABSTRACTS, vol. 125, no. 1, 1 July 1996 Columbus, Ohio, US; abstract no. 11415, XP002080331 & S AIMOTO ET AL.: "Synthesis of phosphorylated calmodulin-binding site of Ca ²⁺ /calmodulin-dependent protein kinase II (CaM KII) by a thioester method " PEPT. CHEM., vol. 32, 1994, pages 197-200, see abstract ---	1-28
A	L E CANNE ET AL.: "Extending the applicability of native chemical ligation" JOURNAL OF THE AMERICAN CHEMICAL SOCIETY., vol. 118, no. 25, 26 June 1996, pages 5891-5896, XP002080328 DC US see the whole document ---	1-28
A	L E CANNE ET AL.: "General method pf thioester resin linkers for use in the solid phase synthesis of peptides alpha-thioacids" TETRAHEDRON LETTERS., vol. 36, no. 8, 20 February 1995, pages 1217-1220, XP002080329 OXFORD GB see the whole document ---	1-28
A	L E CANNE ET AL.: "Synthesis of a versatile purification handle for use with Boc chemistry solid phase peptide synthesis" TETRAHEDRON LETTERS., vol. 38, no. 19, 12 May 1997, pages 3361-3364, XP002080330 OXFORD GB cited in the application see the whole document ---	1-28
	-/-	

INTERNATIONAL SEARCH REPORT

International Application No

PCT/US 98/12278

C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
T	WO 98 28434 A (SCRIPPS) 2 July 1998 see the whole document -----	1-28

INTERNATIONAL SEARCH REPORT

Information on patent family members

Intern/ National Application No

PCT/US 98/12278

Patent document cited in search report	Publication date	Patent family member(s)		Publication date
WO 9634878 A	07-11-1996	AU 2474095 A		21-11-1996
		EP 0832096 A		01-04-1998
WO 9828434 A	02-07-1998	NONE		

THIS PAGE BLANK (USPTO)